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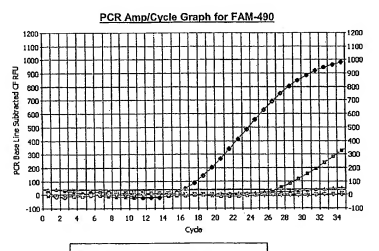
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[Continued on next page]

(54) Title: RAPID DETECTION OF MICROORGANISMS



© zygosaccharomyces bailii
(Lindner) Guilliermond [ATCC
Elndustry-yeast

&B.F.(Mold)

VH₂O(Extraction)

AH₂O(non-extration)

(57) Abstract: Tools and methods for detecting the presence bacteria, yeast and mold in a sample obtained from a food sample are provided. The methods employ a polymerase chain reaction and primer and probe sets that are based on the 16S rRNA and squalene-hopene cyclase genes of Alicyclobacillus and Geobacillus and the 18S rDNA gene of mold and yeast. The present invention also relates to primer and probe sets. Each primer and probe set comprises a forward primer and a reverse primer, both of which are from 15 to 35 nucleotides in length and a probe.





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RAPID DETECTION OF MICROORGANISMS

CROSS REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Provisional Applications No. 60/xxxx, filed October 22, 2003, No. 60/500,736, filed September 05, 2003, and No. 60/430,202, filed December 02, 2002, each of which is incorporated herein by reference in their entirety.

TECHNICAL FIELD OF THE INVENTION

The present invention provides methods and tools for rapidly detecting microorganisms such as molds and fungi, and acid and thermophilic Alicyclobacillus spp and Geobaillus spp. in test samples, particularly food samples.

10 BACKGROUND

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Spoilage of products, particularly food and beverage products, due to contamination with bacteria, yeasts and molds, results in significant financial loss to the food industry. Yeasts and molds are commonly associated with raw materials of foods and are often found in the processing environment. Due to the structural features of both the vegetative cells and spores of fungi, these food contaminants have a good chance of surviving current processing conditions. Yeasts and molds can grow within a wide range of environmental conditions, and therefore the presence in food of even minor amounts of yeast and mold contaminants can cause spoilage during storage.

Like fungi, many bacteria are resitant to processing conditions, and some are resistant even to high acid conditions in food and beverage products. Alicyclobacilli are Gram-positive, spore-forming, aerobic rods classified as thermoacidophiles capable of growing at high temperatures and low pH (1, 2, 3). These bacteria, formerly of the *Bacillus* genus, were assigned into the new genus *Alicyclobacillus* in 1992 (1). Sequence analysis of the 16s rRNA genes proved that three previously classified *Bacillus* thermoacidophiles (*B. acidocaldarius*, *B. acidoterrestris*, and *B. cycloheptanicus*) belong in a group that differs from other closely related Bacilli. Additionally, a key phenotypic variation was found in the membrane composition of these three species. The primary fatty acid component in the membrane was determined to be ω-alicyclic fatty acids, a type of lipid not found in other *Bacillus* species at the time. This evidence initiated the establishment of the *Alicyclobacillus* genus of obligate acidothermophiles, containing *A. acidocaldarius*, *A. acidoterrestris*, and *A. cycloheptanicus*, within the *Bacillus* branch (1). More recently, *A. hesperidum* and *Alicyclobacillus* genomic species 1 and 2 (24, 25), *A. acidiphilus* (22), *A. herbarius* (23), *A. sendaiensis* (26), and *A. pomorum* (27) have been added as new species within the genus *Alicyclobacillus*.

Alicyclobacilli have been an increasingly frequent spoilage problem in the beverage industry, particularly acidic juices, during the last two decades. In 1982, a *Bacillus* sporeformer

(to be later classified as *B. acidoterrestris* and then subsequently *A acidoterrestris*) capable of growing at pH as low as 2.5 was isolated from apple juice (4, 5, 6). In 1994, Splittstoesser et al. discovered the presence of *A. acidoterrestris* in apple juice, further shown by Yamazaki et al. in 1996 (7, 8). Spore germination and growth in orange juice (3) and grapefruit juice (6) was even observed. White grape juice, tomato juice, cranapple juice, and pear juice have also been afflicted with *Alicyclobacillus* spoilage (11).

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While Alicyclobacilli are non-pathogenic, they are a spoilage agent that can drastically affect the quality of acidic fruit juices. Pettipher et al. (1997) reported that guiacol, one of the chemicals responsible for the off-odor and smoky taints characteristic in *Alicyclobacillus*-spoiled juices, can be detected by taste before any visible contamination is seen (3). Therefore, a consumer would generally not be able to identify *Alicyclobacillus*-spoiled juice until it is ingested. In addition to guiacol, 2,6-dibromophenol (2,6-DBP) and 2,6-dichlorophenol (2,6-DCP) were found to contribute to disinfectant taints at detectable levels after as little as one day at 44°C in containers with large headspaces. More realistically, commercially stored shelf stable juices with generally low headspace volume develop these taints within the first month of storage, particularly in warmer climates (10). The presence of these chemicals in *Alicyclobacillus*-spoiled juices significantly reduces the quality of the product, subsequently lowering the consumer image of the brand.

Alicyclobacilli are very heat resistant, growing from pH 2.5-5.5 and 25° C -60° C (6). Beyond growth, cells and spores can survive normal pasteurization procedures, at temperatures up to 97° C (3,6). Fruit juices that are fresh squeezed, pasteurized, or hot-filled are most easily affected by *Alicyclobacillus* spoilage, since ultra high temperature treatment is normally sufficient for killing all microorganisms (3). Since Alicyclobacilli can survive temperatures that exceed industry standard pasteurization specifications, contamination occurring before or during the processing steps can lead to spoilage in the final product that reaches the consumer. Since significant increases in pasteurization temperatures or times ultimately affect product quality and flavor, companies aren't likely to change current procedures.

Early detection, i.e., before products reach the consumer, of the presence of even small amounts of these microbial contaminants in food and beverages is highly desirable in the food industry. Classic culture methods are generally accurate for detecting the presence of microorganisms, but can take up to a week for the results. Previdi et al. (1997) reported a method for detecting A. acidocaldarius in juice products. This method required juices or concentrates to be heat treated and then incubated at 37°C for 7 days, followed by plating on pH 4.0 malt extract agar (13). Pinhatti et al. (1997) tested frozen orange juice concentrate by heat

shocking the samples at 80°C, enriching at 50°C for 24 and 48 h, and finally pour plating in BAM and incubating at 50°C for 24 h (12). Both of these methods of detection provided accurate results, but took from 3-7 days to complete. As with bacteria, it can often take one to two weeks just to grow yeast and mold cells on culture media. In addition, there are so many varieties of molds and yeasts with diverse growth requirements that it is very difficult to find an optimal medium to capture all potential yeast and mold contaminants at the same time. For food industry applications, it is desirable to have a rapid detection system that does not require time consuming culture techniques to detect the presence of microbial contamination of food samples. Accordingly, it is desirable to have a more rapid detection method that can provide results within a few hours, with the same level reliability of culture methods. It is also desirable to have kits that can differentiate between specific types of microbes and which comprise microbe-specific reagents that are useful for conducting rapid sample testing.

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SUMMARY OF THE INVENTION

The present invention provides methods and kits for detecting the presence of Alicyclobacillus spp. and a closely related thermophilic bacterium, Geobacillus, in samples, particularly food samples. In one embodiment the method comprises, collecting bacterial cells in the sample, extracting DNA from the cells, and assaying for the presence of these bacterium species using a PCR technique, preferably real-time PCR, and three signature oligonucleotides (2 primers and a probe) directed to a particular sequence in a target gene encoding either the 16S rRNA or squalene-hopene cyclase (shc). (See the conserved sequences extending from nucleotide position 334 through nucleotide position 485, and from nucleotide position 752 through nucleotide position 813 of the shc gene sequence of Alicyclobacillus shown in Figure 5. Also see the conserved sequences extending from nucleotide position 1327 through nucleotide position 1460 of the 16S rRNA gene sequence of Alicyclobacillus shown in Figure 1.) The presence of multiple Alicyclobacillus spp. and a closely related thermophilic bacterium Geobacillus can be achieved within 3-5 hours using the described sample preparation procedures, and proper combination of the three oligonucleotides as primer-and-probe set in the real-time PCR reaction.

The kits of the present invention comprise at least one forward primer and one reverse primer, with or without a probe for amplifying a sequence of at least 50 consecutive nucleotides within a conserved region of the three Alicyclobacillus spp. shown in Figure 1 (sequences shown in alignment). Figures 2, 3 and 4, respectively, show the full coding sequences for the 16S rRNA genes from the *Alicyclobacillus* strains deposited with the ATCC as 43030, 49025, and 49029. In certain embodiments, the oligonucleotides comprise the entire or a majority of the following

sequences or their reverse complement sequences, as a set or as combination crossing multiple sets, e.g. in certain cases the forward primer of one set can be combined with a reverse primer that is based on the forward primer of another set. Thus the following embodiments can be used in various primer, probe, or primer-probe combinations. Depending on the primers that are combined, the lower oligo may be used as a probe. The sequence of the lower oligo corresponds to the coding sequence of the target region of the gene, and is complementary to the reverse primer in each set. The reverse primers are shown as the reverse complement of the targeted region of the gene. The forward primers correspond to the coding sequence of the target region of the gene.

10	Table I: Signature Oligonucleotides Directed Toward 16S rRNA gene					
		Length	Tm(°C)	GC%		
	Set 1:					
	Forward primer: 5'GAGCCCGCGCGCATTAGC3'	19	68.9	73.7 (SEQ ID NO 1)		
	Probe: 5'GCGACGATGCGTAGCC(G)3'	16	61.8	68.8 (SEQ ID NO 2)		
15	Lower Oligo: 5'CGCAATGGGCGCAAGC3'	16	61.8	68.8 (SEQ ID NO 3)		
•	Reverse primer: 5'GCTTGCGCCCATTGCG3'	16	61.8	61.8 (SEQ ID NO 4)		
	Set 2:					
	Forward primer: 5'GAGCAACGCCGCGTGAGCG3'	19	68.8	73.7 (SEQ ID NO 5)		
20	Probe: 5'CTTCGGGTTGTAAAGC3'	16	54.2	50 (SEQ ID NO 6)		
	Lower Oligo: 5'CGGCTAACTACGTGC3'	15	56.2	60 (SEQ ID NO 7)		
	Reverse primer: 5'GCACGTAGTTAGCCG5'	15	56.2	60 (SEQ ID NO 8)		
	Set 3:					
25	Forward Primer: 5'AGTGCTGGAGAGGCAAGG3'	18	62.2	61.1 (SEQ ID NO 9)		
	Probe: 5'CTGGACAGTGACTGACG3'	17	59.6	58.8 (SEQ ID NO 10)		
	Lower Oligo 5'GCACGAAAGCGTGGGGAGCA	20	66.6	65 (SEQ ID NO 11)		
	Reverse Primer: 5'TGCTCCCCACGCTTTCGTGC5'	20	66.6	65 (SEQ ID NO 12)		
30	Set 4:					
	Forward Primer: 5'GGAGTACGGTCGCAAGACTG3'	20	64.5	60 (SEQ ID NO 13)		
	Probe: 5'CGCACAAGCAGTGGAGC3'	17	62.0	64.7 (SEQ ID NO 14)		
	Lower Oligo: 5'CAGGGCTTGACATC3'	14	52.6	57.1 (SEQ ID NO 15)		
۰.	Reverse Primer: 5'GATGTCAAGCCCTG3'	14	52.6	57.1 (SEQ ID NO 16)		
35	Set 5:					
	Forward primer: 5'GGCGTAAGTCGGAGGAAGG3'	19	64.5	63.2 (SEQ ID NO 17)		
	Probe: 5'ATGTCCTGGGCTACACACG3'	19	62.3	57.9 (SEQ ID NO 18)		
40	Reverse primer: 5'GCCTGCAATCCGAACTACC5'	19	62.3	57.9 (SEQ ID NO 19)		
40	Set CC16S:					
	Forward primer: 5'CGTAGTTCGGATTGCAGGC3'	19	65.6	57.9 (SEQ ID NO 20)		
	Probe: 5'CGGAATTGCTAGTAATCGCG3'	20	57.9	47.4 (SEQ ID NO 21)		
	Lower Oligo: 5'CACGAGAGTCGGCAACAC3'	18	63.3	61.1 (SEQ ID NO 22)		
45	Reverse primer: 5'GTGTTGCCGACTCTCGTG3'	18	62.2	61.1 (SEQ ID NO 23)		

Set 6:

primer: 5'GATGATTGGGGTGAAG3'

16 54.2

50 (SEQ ID NO 24)

Table II: Signature Oligonucleotides Directed Toward squalene-hopene cyclase (shc) gene

These three oligonucleotides were further used as PCR primer pair and DNA probe in real-time PCR detection of *Alicyclobacillus* spp.

Forward Primer: 5' ATGCAGAGYTCGAACG 3' (SEQ ID NO 25)

Probe: 5' 6-FAM d [TCG(A)GAA(G)GACGTCACCGC] BHQ-1 3' (SEQ ID NO 26) Reverse Primer: 5' AAGCTGCCGAARCACTC 3' (Y=C+T; R=A+G (SEQ ID NO 27)

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Table III: The Sequence, GC% and Tm of Primer and probe set candidate 1 for Shc Gene:

Name	Sequence		Tm	GC%
Forward primer	Forward primer TACTGGTGGGGGCCGCT (SEQ ID NO 28)		64.84	70.59
	TACTGGTGGGCGCCGCT (SEQ ID NO 29)	17	64.84	70.59
Probe	ATGGAAGCGGAGTACGTCC (SEQ ID NO 30)	19	62.64	57.9
	ATGGAAGCGGAGTACGTCCT (SEQ ID NO 31)	20	62.45	55
	ATGGAAGCGGAATATGTGC (SEQ ID NO 32)	19	58.32	47.37
	ATGGAAGCGGAATATGTGCT (SEQ ID NO 33)	20	58.35	45
Reverse Primer	CGCGAGGACGCAC (SEQ ID NO 34)	14	62.11	78.57
	CGCGAGGACGCACGTGG (SEQ ID NO 35)	18	69.79	77.78
	CGCGAAGACGGCAC (SEQ ID NO 36)	14	59.16	71.43
	CGCGAAGACGGCACCTGG (SEQ ID NO 37)	18	67.51	72.22

15 Table IV: The Sequence, GC% and Tm of Primer and probe set candidate 2 for Shc General

Name	Sequence	Length	Tm	GC%
Forward primer	CAAAAGGCGCTCGACTG (SEQ ID NO 38)	17	60.02	58.82
	CAAAAGGCGCTCGACTGG (SEQ ID NO 39)	18	62.96	61.1
	CAAAAGGCGCTCGACTGGGTCG (SEQ ID NO 40)	22	68.99	63.64
	CAAAAGTCGCTCGACTG (SEQ ID NO 41)	17	57.61	52.94
	CAAAAGTCGCTCGACTGG (SEQ ID NO 42)	18	60.68	55.56
	CAAAAGTCGCTCGACTGGCTCG (SEQ ID NO 43)	22	67.13	59.09
Probe	GGACGGCGGCTGGGGCGA (SEQ ID NO 44)	18	72.07	83.3
	GGACGCCGCTGGGGCGAGGA (SEQ ID NO 45)	21	75.09	80.9
	GGACGCCGCTGGGGCGAGGACTGCCG (SEQ ID NO 46)	27	80.31	81.4
	GGATGGCGGTTGGGGTGA (SEQ ID NO 47)	18	65.23	66.6
	GGATGGCGGTTGGGGTGAAGA (SEQ ID NO 48)	21	67.28	61.9

	GGATGGCGGTTGGGGTGAAGATTGCCG (SEQ ID NO 49)	27	72.72	62.96
Reverse Primer ^a	TGATGGCGCTCATCGC (SEQ ID NO 50)	16	59.53	62.5
1	TGATGGCGCTCATCGCGGGCGGC (SEQ ID NO 51)	23	74.2	73.91
2	ACCCCGTCGCAGACGGCCTGGGCGC (SEQ ID NO 52)	25	77.7	80
3	ACACCGTCGCAGACCGCCTGGGCGT (SEQ ID NO 53)	25	74.42	72

The present invention also provides methods and kits for detecting the presence of yeast and mold contaminants in samples, particularly in food samples. In one aspect, the method comprises collecting particulate matter, preferably cells and cellular fragments, in the sample, extracting DNA from the particulate matter, and assaying for the presence of yeast DNA in the extracted DNA using a PCR technique using primers that amplify a select conserved region in 18s rDNA of representative yeast species, including *Zygosaccharomyces bailii* (Lindner) Guilliermond strain ATCC 36947 and the other yeast species shown Figure 7. (See conserved sequence extending from nucleotide 81 through nucleotide 225 of the sequence of Z. bali.) Preferably, the method uses real-time PCR, and three signature oligonucleotides (2 primers and a probe) directed to a particular sequence in the gene encoding the yeast 18S rDNA.

In another aspect, the kit of the present invention comprise at least one forward primer and one reverse primer, with or without a probe for amplifying a sequence of at least 50 consecutive nucleotides within the select region of yeast 18s rDNA. In one embodiment, the kit comprises primers and a probe having the following sequences:

Yupreal:

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5' GTGGTGCTAGCATTTGCTG

3' (SEQ ID NO 54)

Ylowreal:

5' GTTAGACTCGCTGGCTCC

3' (SEQ ID NO 55)

Yprobe:

5' TTTCAAGCCGATGGAAGTTTGA(C/G)3' (SEQ ID NO 56)

Another probe that may be used in the present method has the following sequence

20 5' CGGTTTCAAGCCGATGGAAGT 3'. (SEQ ID NO 57)

Yet another set of primers and probe for yeast detection:

Oligo name

Len Pur Scale Sequence (5'-3')

18srRNA-newup-112503-1

25 30 DST 0.05 CCTACTAAATAGGGTGCTAGCATTTGCTGG (SEQ ID NO 58)

18srRNA- newup-112503-2

26 DST 0.05 CTAAATAGGGTGCTAGCATTTGCTGG (SEQ ID NO 59)

18srRNA-probe2

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CGGTTTCAAGCCGATGGAAGTTTGA (SEQ ID NO 60)

In another aspect the present method comprises collecting particulate matter, preferably cells and cellular fragments, in the sample, extracting DNA from the particulate matter, and assaying for the presence of mold DNA in the extracted DNA using a PCR technique using primers that amplify a select conserved region in 18s rDNA of the following representative molds: *Byssochlamys fulva* Olliver et Smith, teleomorph ATCC 24474 and *Penicillium digitatum* Saccardo, anamorph ATCC10030, as shown in the attached alignment. (See the conserved sequence extending from nucleotide 114 through nucleotide 239 of the 18s rDNA sequence of P. digitatum shown in Figure 7.) Preferably, the method uses real-time PCR, and three signature oligonucleotides (2 primers and a probe) directed to a particular sequence in the gene encoding the mold 18s rDNA.

In another aspect, the present the kits of the present invention comprise at least one forward primer and one reverse primer, with or without a probe for amplifying a sequence of at least 50 consecutive nucleotides within the select region of mold 18s rDNA. In one embodiment, the kit comprises primers and a probe having the following sequences:

Mupreal:

5' CCGCTGGCTTCTTAGGG

3' (SEQ ID NO 61)

Mlowreal:

5' AGGGCCAGCGAGTACATCA 3' (SEQ ID NO 62)

Mprobe:

5' CTCAAGCCGATGGAAGTGCG 3' (SEQ ID NO 63)

The invention further provides a method for detecting through real-time PCR using at least one of the nucleic acid primer pairs, and at least one probe, the presence of acidophilic bacterium in a test sample, especially in a food sample. In one embodiment, the acidophilic bacterium detection method includes use of one forward primer directed to the 16S rRNA gene, wherein the primer is selected from the forward primers listed in Table I, one reverse primer directed to the 16S rRNA gene wherein the primer is selected from the reverse primers listed in Table I, and one probe directed to a sequence that is located between the sequences to which the forward and reverse primers are directed, wherein the probe is selected from the group of probles listed in Table I.

In another embodiment, the acidophilic bacterium detection method includes use of one forward primer directed to the squalene-hopene cyclase gene, wherein the forward primer is selected from the group of forward primers listed in Tables II and III, one reverse primer directed to the squalene-hopene cyclase gene wherein the primer is selected from the group of reverse primers listed in Tables II and III, and one probe directed to a sequence that is located between

the sequences to which the forward and reverse primers are directed, wherein the probe is selected from the group of probes listed in Tables II and III.

In yet another embodiment, the acidophilic bacterium detection method includes use of one forward primer directed to the squalene-hopene cyclase gene, wherein the forward primer is selected from the group of forward primers listed in Table IV, one reverse primer directed to the squalene-hopene cyclase gene wherein the primer is selected from the group of reverse primers listed in Table IV, and one probe directed to a sequence that is located between the sequences to which the forward and reverse primers are directed, wherein the probe is selected from the group of probes listed in Table IV.

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In another embodiment, the yeast detection method includes use of one forward primer directed to the 18S rDNA gene, wherein the primer is selected from the group consisting of SEQ ID NO 54 and SEQ ID NO 58, one reverse primer directed to the 18S rDNA gene wherein the primer is selected from the group of consisting of SEQ ID NO 55 and SEQ ID NO 55, and one probe directed to a sequence that is located between the sequences to which the forward and reverse primers are directed, wherein the probe is selected from the group consisting of SEQ ID NO 56, SEQ ID NO 57 and SEQ ID NO 60.

In yet another embodiment, the mold detection method includes use of one forward primer directed to the 18S rDNA gene, wherein the primer corresponds to SEQ ID NO 61, one reverse primer directed to the 18S rDNA gene wherein the primer corresponds to SEQ ID NO 62, and one probe directed to a sequence that is located between the sequences to which the forward and reverse primers are directed, wherein the probe corresponds to SEQ ID NO 63.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows polynucleotide sequence alignment of 16S rRNA gene fragments from three representative strains of *Alicyclobacillus*, specifically, *A. acidocaldarius* ATCC43030, *A. acidoterrestris* ATCC49025, and *A. cycloheptanicus* ATCC49029

Figure 2 shows the 16S rRNA gene coding Sequence for A. cycloheptanicus ATCC49029

Figure 3 shows the 16S rRNA gene coding Sequence for A. acidoterrestris ATCC49025

Figure 4 shows the 16S rRNA gene coding Sequence for A. acidocaldarius ATCC43030

Figure 5: shows the Shc gene sequence alignments for A. cycloheptanicus ATCC49029 and A. acidoterrestris ATCC49025

Figure 6: shows the Shc amino acid sequence alignments for A. cycloheptanicus ATCC49029 and A. acidoterrestris ATCC49025

Figure 7 shows the alignment for the 18s rDNA gene coding Sequence for Zygosaccaromyces, Penecillium digitatum, and Byssochlamys fulva

Figure 8 shows the 16S rRNA gene coding sequence alignments for several strains of for A. cycloheptanicus

Figure 9. shows the results of Real-time PCR detection of A. acidocaldarius (black), A. cycloheptanicus (blue), and A. acidoterrestris (lt. green) using the CC16S specific probe

5 and primer pair.

Figure 10 shows the results of Real-time PCR sensitivity test of A. acidoterrestris using the CC16S primers and probe

Figure 11 shows the results of Real-time PCR sensitivity test of A. acidoterrestris in orange juice.

Figure 12 shows the 18s rDNA gene coding Sequence for Zygosaccaromyces

Figure 13 shows the 18s rDNA gene coding Sequence for Penecillium digitatum

Figure 14 shows the 18s rDNA gene coding Sequence for Byssochlamys fulva

Figure 15 shows the results of a specificity test. © Zygosaccharomyces bailii (Lindner) Guilliermond ATCC 36947;

industry sample yeast. © Byssochlamys fulva Olliver et Smith

ATCC 24474; ∇H_2O control with extraction. $\triangle H_2O$ control without extration.

Figure 16 shows the results of a specificity test with ▼ yeast, •mold and acciobacillus and ▲H₂O

Figure 17 shows the results of a specificity test with \square Z.b(yeast).; \blacktriangle B.F.(mold); \diamondsuit Accidobacillus; \triangledown water; Apple; \triangledown green grape; and \blacksquare Red grape.

Figure 18 shows the results of a specificity test with ■ Z.b.; ▲ B.F.; ♦ Accidobacillus and

Figure 19 shows the results of a specificity test with Gorange1; △ Orange2; ⋄ Orange Juice Supernatant; • Orange Juice pellet; ∘ Yeast; and ▼H₂O

Figure 20 shows the results of a specificity test with !Byssochlamys fulva Olliver et Smith, telomorph ATCC 24474; Penicillium digitatum Saccardo, anamorph ATCC 10030; #Zygosaccharomyces bailii (Lindner) Guillermond, telomorph deposited as Saccharomyces bailii Lindner, telomorph ATCC 36947; %Industry Mold 42; &Industry Mold 41; "Industry Mold 3; "Water (extracted); %water (not extracted)

Figure 21 shows specificity test results with Bussochlamys fulva Olliver et Smith, teleomorph ATCC24474; water and Zygosaccharomyces bailii (Lindner) Guilliermond, telomorph depositied as Saccharomyces bailii Lindner, telomorph ATCC 36947; Acidobacillus acidoterrestris 49025. Figure 22 shows the Alignment^a of 134 bp priming region flanked by CC16S-F (CGTAGTTCGGATTGCAGGC), CC16S-Probe (CGGAATTGCTAGTAATCGC), and CC16S-R (CACGAGAGTCGGCAACAC)^b.

Figure 23 shows the results of Real-time PCR detection of A. acidocaldarius ATCC 43030 (•), A. cycloheptanicus ATCC 49029 (•), and A. acidoterrestris ATCC 49025 (•) using the CC16S primer and probe set.

Figure 24 shows the results of Real-time PCR sensitivity test of A. acidoterrestris ATCC 49025 in saline solution using the CC16S primers and probe

Figure 25 shows the results of Real-time PCR sensitivity test of A. acidoterrestris ATCC 49025 in orange juice, using the CC16S primers and probe.

Figure 26 shows the results of Real-time PCR detection of food-borne microorganisms using the developed primer-and-probe set.

10 Figure 27 shows the resuls f Real-time PCR sensitivity test

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Fig. 28. Real-time PCR detection of A. acidocaldarius ATCC43030 cells in apple juice using shr-specific primer-and-probe set.

DETAILED DESCRIPTION OF THE INVENTION

The methods and kits provided herein enable the rapid and reliable detection of contaminating microorganisms that are found in test samples of products, preferably consumer products, and most preferably food products. The methods are especially suited for the detection of Alicyclobacillus spp. including A. acidocaldarius, A. acidoterrestris, A. cycloheptanicus, A. hesperidum, A. acidiphilus, A. herbarius, A. sendaiensis, and A. pomorum and Geobacillus stearothemophilus, and a variety of yeasts and mold. Other reported methods use conventional PCR (using a pair of oligonucleotides as primers) to detect the presence of Alicyclobacillus spp. (Obara and Niwa, 1998) which usually is associated with the problem of high background with non-specific PCR products.

According to the methods described herein, a sample is obtained from a test material, for example a sample of a fruit juice or other food product. The sample is processed to extract any polynucleotides in the sample, particularly polynucleotides from target organisms that may be present in the material. After extraction and processing according to methods described herein or otherwise known in the art, the sample is treated with reagents that comprise a forward primer oligonucleotide, a reverse primer oligonucleotide, and a labeled oligonucleotide probe, wherein the reagents are targetted for specific regions within the genome of target organisms. The sample is then processed according to PCR amplification methods. The PCR product is first amplified using the primers. Binding of the labeled probe to a target sequence within the PCR product that corresponds with a target region in the genomic DNA of the contaminating bacteria or mold signals the presence of contaminating microorganisms.

Therefore the combination of the three unique sequences and the real-time PCR technology ensured specific and sensitive detection of the presence of the target bacteria. This real-time PCR approach also offers other features such as a) accuracy: more than one probe will be included in the detection system with less possible error; b) flexibility: up to four PCR products can be simultaneously detected so potentially incorporating probes for other spoilage microorganisms into the detection system is expected.

Primer Selection

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Primers are selected within the conserved regions shown in the attached alignment (Figure 1) to amplify a fragment with proper size for optimal detection. One primer is located at each end of the sequence to be amplified. Such primers will normally be between 10 to 35 nucleotides in length and have a preferred length from between 18 to 22 nucleotides. The smallest sequence that can be amplified is approximately 50 nucleotides in length (e.g., a forward and reverse primer, both of 20 nucleotides in length, whose location in the sequences is separated by at least 10 nucleotides). Much longer sequences can be amplified. Preferably, the length of sequence amplified is between 75 and 250 nucleotides in length, and between 75 and 150 for Taqman assay.

One primer is called the "forward primer" and is located at the left end of the region to be amplified. The forward primer is identical in sequence to a region in the top strand of the DNA (when a double-stranded DNA is pictured using the convention where the top strand is shown with polarity in the 5' to 3' direction). The sequence of the forward primer is such that it hybridizes to the strand of the DNA which is complementary to the top strand of DNA.

The other primer is called the "reverse primer" and is located at the right end of the region to be amplified. The sequence of the reverse primer is such that it is complementary in sequence to, i.e., it is the reverse complement of a sequence in, a region in the top strand of the DNA. The reverse primer hybridizes to the top strand of the DNA.

PCR primers should also be chosen subject to a number of other conditions. PCR primers should be long enough (preferably 10 to 30 nucleotides in length) to minimize hybridization to greater than one region in the template. Primers with long runs of a single base should be avoided, if possible. Primers should preferably have a percent G+C content of between 40 and 60%. If possible, the percent G+C content of the 3' end of the primer should be higher than the percent G+C content of the 5' end of the primer. Primers should not contain sequences that can hybridize to another sequence within the primer (i.e., palindromes). Two primers used in the same PCR reaction should not be able to hybridize to one another. Although PCR primers are preferably chosen subject to the recommendations above, it is not necessary that the primers

conform to these conditions. Other primers may work, but have a lower chance of yielding good results.

PCR primers that can be used to amplify DNA within a given sequence can be chosen using one of a number of computer programs that are available. Such programs choose primers that are optimum for amplification of a given sequence (i.e., such programs choose primers subject to the conditions stated above, plus other conditions that may maximize the functionality of PCR primers). One computer program is the Genetics Computer Group (GCG recently became Accelrys) analysis package which has a routine for selection of PCR primers. There are also several web sites that can be used to select optimal PCR primers to amplify an input sequence. One such web site is http://alces.med.umn.edu/rawprimer.html. Another such web site is http://www-genome.wi.mit.edu/cgi-bin/primer/primer3_www.cgi.

Making the Oligonucleotide Primers and Probes

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The oligonucleotide primers and probes disclosed in this application can be made in a number of ways. One way to make these oligonucleotides is to synthesize them using a commercially-available nucleic acid synthesizer. A variety of such synthesizers exists and is well known to those skilled in the art. Many such synthesizers use phosphoramidite chemistry, although other chemistries can be used. Phosphoramidite chemistry utilizes DNA phosphoramidite nucleosides, commonly called monomers, to synthesize the DNA chain or oligonucleotide. Such monomers are modified with a dimethoxytrityl (DMT) protecting group on the 5'-end, a b-cyanoethyl protected 3'-phosphite group, and may also include additional modifiers that serve to protect reactive primary amines in the heterocyclic ring structure (to prevent branching or other undesirable side reactions from occurring during synthesis).

To make an oligonucleotide of a specific sequence, phosphoramidite nucleosides are added one-by-one in the 3'-5' direction of the oligonucleotide, starting with a column containing the 3' nucleoside temporarily immobilized on a solid support. Synthesis initiates with cleavage of the 5'-trityl group of the immobilized 3' nucleoside by brief treatment with acid [dichloroacetic acid (DCA) or trichloroacetic acid (TCA) in dichloromethane (DCM)] to yield a reactive 5'-hydroxyl group. The next monomer, activated by tetrazole, is coupled to the available 5'-hydroxyl and the resulting phosphite linkage is oxidized to phosphate by treatment with iodine (in a THF/pyridine/H₂O solution). The above describes the addition of one base to the oligonucleotide. Additional cycles are performed for each base that is added. The final oligonucleotide added does not have a 5' phosphate. When synthesis is complete, the oligonucleotide is released from the support by ammonium hydroxide and deprotected (removal of blocking groups on nucleotides).

Normally, oligonucleotides of up to 150-180 bases long can be efficiently synthesized by this method using a nucleic acid synthesizer. To make oligonucleotide that are longer than 100 bases, two single-stranded oligonucleotides, that are partially complementary along their length, can be synthesized, annealed to one another to form a duplex, and then ligated into a plasmid vector. Once a plasmid containing the ligated duplexes has been formed, additional oligonucleotide duplexes can be ligated into the plasmid, adjacent to the previously ligated duplexes, to form longer sequences. It is also possible to sequentially ligate oligonucleotide duplexes to each other, to form a long, specific sequence, and then clone the single long sequence into a plasmid vector.

Sample preparation flow chart for bacteria detection

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Collect cells by centrifugation or membrane filtration

Lyse Cells using standard techiniques

DNA extraction using standard techniques

Analysis (Real-time PCR)

Sample preparation flow chart for fungi (yeast and mold) detection

Collect cells and cell fragments by centrifugation or membrane filtration

Lyse Cells using standard techniques

Extract DNA using standard techniques

Analysis (Real-time PCR)

Isolation of DNA from Samples

DNA is isolated or extracted from the microorganism cells contained within the test sample. For example, DNA extraction may be performed using any of numerous commercially available kits for such purpose. One such kit, called IsoCode, is available from Schleicher and Schuell of Keene, New Hampshire. The IsoCode kit contains paper filters onto which cells are applied. Through treatment of the paper filters as described by the manufacturer, most cellular components remain in the paper filter and DNA is released into an aqueous solution. The DNA in the solution can then be added to various enzymatic amplification reactions, as are discussed below.

Other commercially available kits exist for extraction of DNA from cells. Commercial kits do not have to be used, however, in order to obtain satisfactory DNA. Standard methods, well known to those skilled in the art, have been published in the scientific literature. Such methods commonly involve lysis of cells and removal of cellular components other than nucleic acids by precipitation or by extraction with organic solvents. Enzymatic treatment with proteases and ribonucleases can be used to remove proteins and RNA, respectively. DNA is then commonly precipitated from the sample using alcohol.

Real-Time PCR

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A variety of methods can be used to determine if a PCR product has been produced. One way to determine if a PCR product has been produced in the reaction is to analyze a portion of the PCR reaction by agarose gel electrophoresis. For example, a horizontal agarose gel of from 0.6 to 2.0% agarose is made and a portion of the PCR reaction mixture is electrophoresed through the agarose gel. After electrophoresis, the gel is stained with ethidium bromide. PCR products are visible when the gel is viewed during illumination with ultraviolet light. By comparison to standardized size markers, it is determined if the PCR product is of the correct expected size.

The PCR procedure preferably is done in such a way that the amount of PCR products can be quantified. Such "quantitative PCR" procedures normally involve comparisons of the amount of PCR product produced in different PCR reactions. A number of such quantitative PCR procedures, and variations thereof, are well known to those skilled in the art. One inherent property of such procedures, however, is the ability to determine relative amounts of a sequence of interest within the template that is amplified in the PCR reaction.

One particularly preferred method of quantitative PCR used to quantify copy numbers of sequences within the PCR template is a modification of the standard PCR called "real-time PCR." Real-time PCR utilizes a thermal cycler (i.e., an instrument that provides the temperature changes necessary for the PCR reaction to occur) that incorporates a fluorimeter (i.e. an instrument that measures fluorescence). In one type of real-time PCR, the reaction mixture also contains a reagent whose incorporation into a PCR product can be quantified and whose quantification is indicative of copy number of that sequence in the template. One such reagent is a fluorescent dye, called SYBR Green I (Molecular Probes, Inc.; Eugene, Oregon) that preferentially binds double-stranded DNA and whose fluorescence is greatly enhanced by binding of double-stranded DNA. When a PCR reaction is performed in the presence of SYBR Green I, resulting DNA products bind SYBR Green I and fluoresce. The fluorescence is detected

and quantified by the fluorimeter. Such technique is particularly useful for quantification of the amount of template in a PCR reaction.

A preferred variation of real-time PCR is TaqMan® (Applied Biosystems) PCR. The basis for this method is to continuously measure PCR product accumulation using a dual-labeled flourogenic oligonucleotide probe called a TaqMan® probe. The "probe" is added to and used in the PCR reaction in addition to the two primers. This probe is composed of a short (ca. 15-30 bases) oligodeoxynucleotide sequence that hybridizes to one of the strands that are made during the PCR reaction. That is, the oligonucleotide probe sequence is homologous to an internal target sequence present in the PCR amplicon. The probe is labeled or tagged with two different flourescent dyes. On the 5' terminus is a "reporter dye" and on the 3' terminus is a "quenching dye." One reporter dye that is used is called 6-carboxy fluorescein (FAM). One quenching dye that is used is called 6-carboxy tetramethyl-rhodamine (TAMRA). When the probe is intact, energy transfer occurs between the two fluorochromes and emission from the reporter is quenched by the quencher, resulting in low, background fluorescence. During the extension phase of PCR, the probe is cleaved by the 5' nuclease activity of Taq polymerase, thereby releasing the reporter from the oligonucleotide-quencher and producing an increase in reporter emission intensity. During the entire amplification process the light emission increases exponentially.

Because the detection in Taqman assay is based on complementary binding of the third oligonucleotide probe to the amplified PCR products, it can significantly minimize false positive results due to the detection of non-specific amplification and primer dimers in conventional PCR and other non-specific real-time PCR product detection approaches such as using SYBR Green or EtBr. However, the determination of proper primer and probe set needs more specified skills so that they will fit the product amplification and signal detection requirements.

Examples of primers and probes that are particularly useful in this procedure are listed above.

Fluorescence Detection

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One example of an instrument that can be used to detect the fluorescence is an ABI Prism 7700, which uses fiber optic systems that connect to each well in a 96-well PCR tray format. The laser light source excites each well and a CCD camera measures the fluorescence spectrum and intensity from each well to generate real-time data during PCR amplification. The ABI 7700 Prism software examines the fluorescence intensity of reporter and quencher dyes and calculates the increase in normalized reporter emission intensity over the course of the amplification. The results are then plotted versus time, represented by cycle number, to produce a continuous

measure of PCR amplification. To provide precise quantification of initial target in each PCR reaction, the amplification plot is examined at a point during the early log phase of product accumulation. This is accomplished by assigning a fluorescence threshold above background and determining the time point at which each sample's amplification plot reaches the threshold (defined as the threshold cycle number or CT). Differences in threshold cycle number are used to quantify the relative amount of PCR target contained within each tube.

Detecting fungi in samples

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Oligonucleotide primer and probe development for detecting yeast

We have cloned and sequenced the 18s rDNA gene fragments from representative yeast Zygosaccharomyces bailii (Lindner) Guilliermond strain ATCC 36947. We then compared our sequences against other published 18S rDNA sequences from molds, yeasts, and common eukarytic foods. We have also compared other target sequences including h1, h2, 23S rDNA, spacer sequence between 18S and 23S rDNA gene. We have developed primer-and-probe sequences that can detect the presence of generally all yeasts without cross-reacting with foods, molds or other bacteria. The aligned sequences of the 18S rDNA sequences of these yeast species are shown in Figure 17. Figures 12, 13 and 14 show the full coding sequences for the genes corresponding to the alignments shown in Figure 17.

Specificity Testing

Using the primer pair-and-probe set, all yeasts were tested positive in real-time PCR (Figures 15-19), while no cross-reaction was detected in other commonly found foodborne microorganisms and food items (Figures 15-19). Further specificity study revealed no combination of the above three oligonucleotides in other microorganisms after blast searching the nucleotide sequence database in the GenBank.

Oligonucleotide primer and probe development for detecting mold

We have cloned and sequenced the 18s rDNA gene fragments of representative molds of 25 food industry concerns, Byssochlamys fulva Olliver et Smith, teleomorph ATCC 24474 and Penicillium digitatum Saccardo, anamorph ATCC10030. Coloning primer up:TGCATGGCCGTTCTTAGTTGG(Z.B. code 64-75) (B.F. 667-688) (P.D. 674-695) down: GTGTGTACAAAGGGCAGGG(Z.B. 417-237) (B.F. 1011-1031) (P.D. 1029-1049). We 30 then compared our sequences against other published 18S rDNA sequences from molds, yeasts, and common eukarytic foods. We have also compared other target sequences including h1, h2, 23S rDNA, spacer sequence between 18S and 23S rDNA gene. We have developed primer and probe sequences that can detect the presence of generally all mold without cross-reacting with foods, yeast or bacteria.

Specificity test

Using primer pair-and-probe set, all yeasts were tested positive in real-time PCR (Figures 20 and 21), while no cross-reaction was detected in other commonly found foodborne microorganisms and food items (Figures 20 and 21). Further specificity study revealed no combination of the above three oligonucleotides in other microorganisms after blast searching the nucleotide sequence database in the GenBank.

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EXAMPLES

Example 1:

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In this study, the 16s rDNA sequences of A. acidocaldarius, A. cycloheptanicus, and A. acidoterrestris were used as models for the development of specific primers and a flourogenic 10 probe to be used in a real-time PCR assay. 16s rDNA was isolated from ATTC strains 43030, 49025, and 49029, then cloned into vectors, transformed into competent cells, and purified for sequencing. Following sequencing, the 16s rDNA sequences of the three strains were analyzed for the development of oligonucleotide primers and a flourescent probe. These primers and probe were used in a real-time PCR detection system where specificity and sensitivity tests were 15 performed in media as well as beverage systems. This rapid detection system is unique because it can specifically detect not only the three original Alicyclobacillus species, but also detects newer species of Alicyclobacillus because of the genus-level 16s rDNA conservation of the priming sequences. This system can greatly benefit the food industry, particularly the beverage industry, by detecting the presence of Alicyclobacillus within hours, before the product ever 20 reaches the consumer, saving not only time and money, but maintaining brand image and quality. Materials and Methods

Bacterial strains and culture conditions. A. acidocaldarius strain ATTC 43030 was grown on

ATCC 573 medium, consisting of 1.3g (NH₄)₂SO₄, 0.37g KH₂PO₄, 0.25g MgSO₄·7H₂O, 0.07g

CaCl₂·2H₂O, 1.0g glucose, 1.0g yeast extract, and 1.0 L distilled H₂O. Solution pH was adjusted

to 4.0 using H₂SO₄ and autoclaved at 121°C for 15 minutes. A. acidoterrestris strain ATTC 49025 and A. cycloheptanicus strain ATCC 49029 were grown on BAM-SM ATCC 1656

 $medium\ consisting\ of\ 0.25g\ CaCl_2\cdot 2H_2O,\ 0.5g\ MgSO_4\cdot 7H_2O,\ 0.2g\ (NH_4)_2SO_4,\ 3.0g\ KH_2PO_4,\ 6.0g$

yeast extract, 5.0g glucose, 1.0mL trace elements (0.66g CaCl₂·2H₂O, 0.18g ZnSO₄·7H₂O, 0.16g

CuSO₄·5H₂O, 0.15g MnSO₄·4H₂O, 0.18g CoCl₂·6H₂O, 0.10g H₃BO₃, 0.30g Na₂MoO₄·2H₂O,

1.0L distilled H₂O_{),} and 1.0L distilled H₂O. Solution pH was adjusted to 4.0 using H₂SO₄ and autoclaved at 121°C for 15 minutes. Stock cultures of all strains were stored in their respective

media plus 40% glycerol and kept at -80°C.

Isolation of genomic DNA and amplification of 16s rDNA. DNA was isolated from 2% cultures of A. acidoterrestris strain ATTC 49025, A. cycloheptanicus strain ATCC 49029 A. acidocaldarius strain ATTC 43030 in respective media. Cultures were grown for 24 hours at

47°C. Genomic DNA was extracted from each strain using the Qiagen DNeasy Tissue Kit (Qiagen, Valencia, CA). The included protocol was followed, except the elution was repeated once with 100μl of buffer AE. An approximately 1,500 bp region of the 16s rDNA was amplified from the genomic DNA using primers 8F and 1492R (15) with PCR performed on the Bio-Rad iCycler Thermal Cycler (Bio-Rad Laboratories, Hercules, California). A 50μl reaction mixture was used, containing 0.5μl of primer 8F, 0.5μl of primer 1492R, 1.0μl of genomic DNA, 37μl of sterile H₂O, 3μl of 50mM MgCl₂, 2μl of a 10mM dNTP mixture, and 1.0μl Taq polymerase (Invitrogen, Carlsbad, CA). Amplification conditions included 30 cycles of 95°C for 2 min, 42°C for 30 s, and 72°C for 4 min, with a final chain elongation for 20 min (15). PCR products were confirmed after 20 min of gel electrophoresis on 0.9% agarose gel at 100 volts, followed by 10 min of ethidium bromide staining for visualization.

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Cloning and transformation of 16s rDNA gene. PCR products were purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA). The protocol was followed as specified by the manufacturer, except 30µl of sterile H₂O was used in place of 50µl of buffer EB for a single elution. Purified PCR products were then cloned into pCR 2.1 vectors using the TA Cloning kit (Invitrogen, Carlsbad, CA). A 10 µl ligation reaction for each PCR product was prepared as follows: 5µl sterile H₂O, 1µl pCR 2.1 vector, and 2µl PCR product were mixed together and incubated at 65°C for 5 min, followed by 10 min of incubation on ice. 1µl 10X ligation buffer and 1µl T4 DNA ligase were then added to the mixture, followed by overnight incubation at Transformation was then performed, beginning with centrifugation of the ligation reactions. Reactions were stored on ice while 50µl of One Shot competent Escherichia coli cells were thawed for each transfer. 5µl of each ligation reaction was added to a vial of One Shot cells and mixed gently, followed by incubation for 30 min on ice. Reactions were then heat shocked for 30 s at 42°C, and then placed on ice. 200µl of SOC medium was added to each tube and then shook at 200 rpm for one hour at 37°C. The whole vial of cells was then spread onto LB agar plates containing X-Gal (20mg/ml) and incubated at 37°C overnight. Plates were stored at 4°C following incubation.

Sequencing of 16s rDNA gene. Plates were observed for transformed (white) colonies. Five transformed colonies from each plate were selected using a sterile toothpick, then dipped into a microfuge tube containing 100μl of sterile H2O, and also spread on an LB agar plate. The stick was then placed into a tube containing 2ml of LB broth and ampicillin (50mg/ml). Plates were incubated at 37°C overnight. LB tubes were shaken at 100 rpm at 37°C overnight. Microfuge tubes were incubated at 100°C for 10 min, followed by PCR to check for successful

transformation. Standard 3-step PCR (CYCLES) was run with a 50µl reaction mixture containing 0.5µl of primer M13F, 0.5µl of primer M13R, 1.0µl of transformed DNA, 37µl of sterile H₂O, 3µl of 50mM MgCl₂, 2µl of a 10mM dNTP mixture, and 1.0µl Taq polymerase (Invitrogen, Carslbad, CA). PCR products were analyzed by gel electrophoresis. LB tubes were centrifuged for 10 min at 6000 rpm after overnight incubation and used in the QIAprep Spin Miniprep kit (Qiagen, Valencia, CA) following the manufacturer's protocol. 5µl of product was set aside for PCR, and the rest of the miniprep yield was sent to be sequenced. Sequence data was entered into the NCBI BLAST network to search for similar sequences. Cloned sequences from ATCC strains 49025, 49029, and 43030 matched multiple 16s rDNA sequences from Alicyclobacillus species on the BLAST network.

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Real-time Taqman PCR conditions. Fifty microliter reaction mixtures containing 0.5μl of a 100μM solution of CC16S-F primer, 0.5μl of a 100μM solution of CC16S-Probe, 33.3μl of sterile H₂O, 5.0μl of genomic DNA, 5μl of 10X reaction buffer, 3μl of MgCl₂, 2μl of dNTP's, and 0.2μl of Taq polymerase (Invitrogen, Carlsbad, CA) were used for specificity tests. For sensitivity assays, the following 50μl reaction mixtures were used: 25μl of 2X iQ Supermix, containing 100 mM KCl, 40 mM Tris-HCl, pH 8.4, 0.4 mM each dNTP, 50 U/ml iTaq DNA polymerase, 6 mM MgCl₂, and stabilizers (Bio-Rad, Hercules, CA), 0.5μl of 100μM stock CC16S-F primer, 0.5μl of 100μM stock CC16S-R primer, 0.5μl of 100μM stock CC16S-Probe, 5.0μl of genomic DNA, and 18.5μl of sterile H₂O. Real-time PCR was performed using the iCycler iQ Real-Time PCR Detection System (Bio-Rad, Hercules, CA). PCR conditions were as follows: 35-40 cycles of 95°C denaturation for 30 s and 55°C annealing for 30 s. The optical module was set to capture light during the annealing step. Results were analyzed using the iCycler iQ Optical System Software Version 3.0a (Bio-Rad, Hercules, CA).

Primer and probe design. Sequence alignments of the 16s rDNA sequences for strains 49025, 29029, and 43030 were constructed with ClustalV using MegAlign 5.01 (DNASTAR, Madison, WI). A sequence alignment of the 16S rDNA sequences was then performed for the following organisms: sequenced Alicyclobacillus strains ATCC 49025, 49029, and 43030, A. acidoterrestris strain DSM 3923 (AB042058), A. cycloheptanicus strain DSM 4006 (AB042059), A. acidocaldarius strain DSM 454 (AB059664), Geobacillus subterraneus strain K (AF276307), Sulfobacillus disulfidooxidans SD-11 (U34974), B. thermoleovorans strain ATCC 43513 (M77488), and Clostridium elmenteitii isolate E2SE1-B (AJ271453). The alignment was constructed with ClustalV using MegAlign 5.01 (DNASTAR, Madison, WI). Aligned regions were carefully scanned by eye to find areas of perfect identity within the representative

Alicyclobacillus species in order to create PCR priming regions. The following criteria were used for primer and probe selection: (1) 100% identity between representative sequences, (2) priming region of less than 200 bp, $(3)T_m$ greater than 55°C, (4) C or G in the terminal positions of both 5' and 3' ends, (5) greater than 45% C+G content, and (6) no visual hairpin loops or secondary structures, confirmed using the Oligo Toolkit (Qiagen, Valencia, CA) (22).

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Specificity and sensitivity tests. Assays were performed using the aforementioned PCR conditions to test for specificity of the system for Alicyclobacillus spp. and any cross-reactions with other common food-borne microorganisms. Genomic DNA was extracted from broth cultures of 2% A. acidoterrestris, A. acidocaldarius, and A. cycloheptanicus grown for 48 h at 47°C using the previously discussed DNA extraction protocol. In addition, genomic DNA was extracted from Escherichia coli DH-5\alpha, Lactococcus lactis subsp. lactis, Geobacillus stearothermophilus ATCC 10149 and Pseudomonas putida 49L/51 to test specificity of the primers and probe.

Assays for the sensitivity of the real-time PCR assay for detection of Alicyclobacillus were performed using tenfold serial dilutions of 10° to 10⁻⁸ of A. acidoterrestris in a 10 ml 15 solution of 0.85% NaCl. Two percent cultures were initially grown for 48 h at 47°C in order to. obtain an OD₆₀₀ range between 0.400 and 0.800. After dilution, cells from 1ml of each sample were collected by centrifugation at 12,000 rpm for 10 minutes for DNA extraction. Fifty microliter (50µl) reaction mixtures containing 0.5µl of CC16S-F primer, 0.5µl CC16S-R primer, 20 0.5µl CC16S-Probe, 33.3µl of sterile H₂O, 5.0µl of genomic DNA, 5µl of 10X reaction buffer, 3μl of MgCl₂, 2μl of dNTP's, and 0.2μl of Taq polymerase (Invitrogen, Carlsbad, CA) were used for each strain, as described above. Real-time PCR was carried out with the following cycling conditions: 35-40 cycles of 95°C and 55°C, for 30 s each. After amplification, results were analyzed using the iCycler iQ Optical System Software Version 3.0a (Bio-Rad, Hercules, CA). A range of dilutions between 10⁻³ and 10⁻⁷ were plated on BBL Orange Serum Agar (Difco. 25 Detroit) for colony counting. Plates were incubated at 47°C for 48h. Additionally, sensitivity tests were performed in the same manner using apple and orange juice. Also, 1ml of culture was spiked in 9ml of Powerade sports drinks and Minute Maid Lemonade to check for any inhibitory characteristics these drinks may display in a PCR assay.

Amplification, cloning, transformation, and sequencing of 16s rDNA gene. PCR was used to successfully amplify regions of 16s rDNA from A. acidoterrestrs, A. acidocaldarius, and A. cycloheptanicus using the 8F and 1492R primers. The Invitrogen TA cloning kit was used to insert the amplified 16s rDNA segment of each strain into pCR 2.1 vectors, and subsequently transformed into E. coli competent cells. Purified samples were then sent to the Plant-Microbe

Genomic Facility at the Ohio State University and sequenced using an ABI PRISM 3700 DNA Analyzer (Applied Biosystems, Foster City, CA).

TABLE V. Oligonucleotide data for Alicyclobacillus spp. CC16S probe and primers.

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Name	Sequence	Length	T_m	G+C content
CC16S-F	CGTAGTTCGGATTGCAGGC	19 bp	65.6°C	57.9%
CC16S-R	GTGTTGCCGACTCTCGTG	18 bp	63.3°C	61.1%
CC16S-Probe	CGGAATTGCTAGTAATCGC	19 bp	57.9°C	47.4%

Development of CC16S primers and probe. Sequence data obtained from the Plant-Microbe Genomics Facility was compiled and entered into the NCBI BLAST network to check sequence integrity. Sequence data for each strain corroborated with respective sequence data in the GenBank. The 16S rDNA sequences from the three sequenced strains, as well as from A. acidoterrestris strain DSM 3923 (AB042058), A. cycloheptanicus strain DSM 4006 (AB042059), and A. acidocaldarius strain DSM 454 (AB059664) were used as positive controls in the alignment to determine a suitable priming region. B. thermoleovorans strain ATCC 43513 (M77488) and Clostridium elmenteitii isolate E2SE1-B (AJ271453) were used as negative controls in the alignment. In addition, closely related Geobacillus subterraneus strain K (AF276307) and Sulfobacillus disulfidooxidans SD-11 (U34974) were added to the alignment. Using the criteria described in the methodology, a forward and reverse primer and fluorogenic probe were derived, named CC16S-F, CC16S-R, and CC16S-Probe respectively. The sequences for the oligonucleotides are shown in Table V. This oligonucleotide set will amplify a 134 bp segment of the 16S rDNA. The alignment of the 134 bp priming region is shown in Figure 22, with the selected primer and probe oligonucleotide sequences boxed around the Alicyclobacillus strains. These sequences were entered into the BLAST search network in order to discover identities with other unrelated organisms to ensure their specificity for Alicyclobacillus. Results show that the priming sequences are specific for 16S rDNA sequences of the three Alicyclobacillus species sequenced. In addition, the priming sequences also match the newly discovered species A. hesperidum, A. herbarius, A. acidiphilus, and A. sendaiensis. Also, it was found after alignment and BLAST searches that the priming region was highly similar to the members of the Geobacillus and Sulfobacillus genera, two closely related groups. Primers CC16S-F and CC16S-R were ordered from Sigma-Genosys (The Woodlands, TX), and the CC16S-Probe was ordered from Biosearch Technologies (Novato, CA). CC16S-Probe was labeled with the reporter dye Quasar 670 on the 5' end, and quencher dye BHQ-2 on the 3' end.

Real-time PCR specificity assay. Real-Time PCR is a new method has been developed to overcome the problems of standard PCR while increasing sensitivity and allowing for nearly instantaneous results. Real-time PCR adds an optical module and a fluorogenic probe to a standard PCR assay, while including computer-based data analysis software for real-time monitoring. Real-time PCR eliminates the need for post-amplification analysis and is not affected by non-specific amplification. The optical module attached to the thermal cycler detects a flourescent signal that is emitted from the labeled probe at each cycle during the annealing stage. The amount of emission is recorded by computer software and plotted as an exponential curve, displaying the cycle at which a significant amount of amplification takes place.

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The flourescent reporter dye is held on the 5' end of an oligonucleotide probe, with a quenching dye on the 3' end to capture flourescence not related to amplification. When the probe anneals within the primed region, the 5' exonuclease activity of the polymerase in the reaction system cleaves the probe, inhibiting the quencher dye and increasing the emitted flourescence from the 5' reporter dye (21).

A real-time PCR assay was developed to test the specificity of the primers and probe for A. acidoterrestris, A. acidocaldarius, and A. cycloheptanicus. The assay also included E. coli DH-5 α , L. lactis subsp. lactis, and P. putida to test for any unwanted cross-reactions with common foodborne microorganisms. In addition, Geobacillus stearothermophilus ATCC 10149 was included in the assay since it is a closely related thermophile of the Bacillus subfamilies. Assays were performed in triplicate, and results analyzed using the iCycler iQ Optical System Software. The results show that the reaction is specific for the three Alicyclobacillus while not reacting with E. coli DH-5 α , E. lactis subsp. lactis, or E0 putida. However, E0 stearothermophilus had a positive reaction within the system.

Real-time PCR sensitivity assay and limit of detection. After establishing system specificity, sensitivity of detection was determined. In order to accomplish this, tenfold serial dilutions in a 0.85% NaCl solution were made using A. acidoterrestris ATCC 49025 cultures. Real-time PCR assays were run in triplicate and results were analyzed using the iCycler iQ Optical System Software. A typical result is shown in Figure 23. Quantification of the lowest detection level was performed through colony counting of plated dilutions used in the PCR. Colonies were counted on OSA plates and then averaged. The CFU/ml was calculated, and cell counts were determined for the lowest positive curve by multiplying the CFU/ml by the dilution factor of the curve. Data for cell counts and detection limits is presented in Table VI. In Figure 24, the lowest accurate curve presented is from a 10⁻⁵ dilution, which is equivalent to 160 CFU/ml by plate

count. Sensitivity tests were performed in triplicate, with the limit of detection ranging between 66 and 160 cells. The mean detection limit is 103 cells.

TABLE VI. A. acidoterrestris cell counts and corresponding detection limits for sensitivity tests performed in saline solution and orange juice.

Replicate	Media	Mean number of colonies ^a	Mean cell count per replicate (CFU/ml)	Minimum PCR detection level per replicate ^c	Mean PCR detection level for trial set ^d
1	Saline	8	8.3 x 10 ⁶ b	8.3 x 10 ¹	Saline solution
2	Saline	160	1.60×10^7	1.60×10^2	1.03×10^{2}
3	Saline	66	6.6×10^6	6.6 x 10 ¹	
1	Orange Juice	21	2.1×10^7	2.1 x 10 ¹	Orange juice
2	Orange Juice	63	6.3×10^7	6.3×10^{1}	5.36 x 10 ¹
3	Orange Juice	76	7.6×10^7	7.6 x 10 ¹	

^a Diluted samples of A. acidoterrestris in respective media were plated on replicate plates of BBL Orange Serum Agar (Difco, Detroit), and colony counts and averages were obtained after 48h at 47°C.

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The detection limit of the *Alicyclobacillus* real-time PCR rapid screening system was also established in beverages using orange juice as a diluent. Serial dilutions were performed as previously described with juice in place of 0.85% NaCl. Juice samples were initially run in parallel with samples in 0.85% NaCl, and C_T values and curve intensities were found to be comparable in both systems. Results for the assay in orange juice are shown in Figure 25. Colony counting was performed on plated dilutions used in the PCR in order to determine cell counts at the minimum detection level. Data for cell counts and detection limits is presented in Table VI. In Figure 25, the lowest accurate curve presented is from a 10^{-6} dilution, which is equivalent to 63 CFU/ml. Sensitivity tests were performed in triplicate, with the limit of detection ranging between 21 and 76 cells. The mean detection limit is 54 cells.

The efficiency of the system has also been tested in other beverages including apple juice, three sports drinks and Lemonade purchased from local grocery stores. These beverages were spiked with A. acidoterrestris cultures followed by cell collection, DNA extraction and real-time

^b Calculation is estimated because no plates with between 20 and 200 colonies were available.

^c Minimum detection level is calculated by multiplying the mean cell count per replicate by the dilution level of lowest positive real-time PCR detection curve from the corresponding amplification run.

d This is the calculated average detection limit for repeated real-time PCR trials in each type of media.

PCR detection. In all these cases, expected PCR amplification results were obtained indicating no particular inhibition by the ingredients from these tested beverages.

Discussion

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A specific and sensitive real-time PCR-based rapid detection system for Alicyclobacillus has been developed. In the past, PCR based assays have been used to detect microorganisms in different environments (16, 2, 17, 18, 19, 20, 28). More recently, the use of real-time PCR has been a favorable alternative to standard PCR based assays due to the increased speed and sensitivity of the results, the ability to quantify detection levels, and the elimination of postamplification analysis (21). The present method was developed by targeting the 16s rDNA gene of Alicyclobacilli, using A. acidoterrestris, A. acidocaldarius, and A. cycloheptanicus as models for primer and probe development. However, the developed primers and probe could also be beneficial in detecting newly classified members of Alicyclobacillus, due to high sequence identity as shown by the BLAST data. This real-time PCR assay is an improvement over traditional culture methods of detection and PCR based detection systems. Culture methods can take between three and seven days for results to be available (12, 13). While accurate, the time frame is much too long for practical industry implementation. PCR assays provide much quicker results, but false positives can be easily detected (21), and gel electrophoresis analysis must be performed after amplification. Real-time PCR assays can be readily implemented in the industry because of the real-time results. Samples can be taken from the floor as they are produced and the presence of Alicyclobacilli can be detected within 3 hours.

In this study, the developed primers and probes were able to specifically detect A. acidocaldarius, and A. cycloheptanicus without cross-reaction with other common foodborne microorganisms. In addition, the system could also detect the presence of G. stearothermophilus.

25 Example 2:

A real-time PCR based rapid system was developed for detecting spoilage Alicyclobacillus spp. in foods. A common gene of Alicyclobacillus spp. encoding squalene-hopene cyclase, a key enzyme involved in hopanoid biosynthesis, was targeted for specific primers and probe development. Using the combination of the primers and probe, specific detection of the presence of representative strains from Alicyclobacillus spp. was achieved in the Taqman-based real-time PCR assay without cross-reacting with other food-borne bacteria. The presence of around 100 cells in collected samples can be detected within several hours.

Food spoilage causes significant financial loss to the industry. Every year, about 10% of our food supplies are lost due to spoilage and a significant portion of the problem is because of

the presence of spoilage microbial agents, particularly molds, yeasts, and bacteria capable of surviving moderate heat- and acidic-treatments. Due to the limitation of applying extreme processing conditions, which can significantly alter the physiochemical properties and nutritional values of many food products, proper detection screening for the presence of microbial spoilage agents in food becomes a prior choice for quality control in the food industry. However, conventional industry practices for microbial detection from plate counting to biochemical analysis take anywhere from 48 hours to a couple of weeks. These methods are especially unsuitable for products with limited shelf life such as fruit juices. Novel detection approaches enabling rapid and specific detection of spoilage microorganisms within hours are preferred.

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While the polymerase chain reaction (PCR) has been used extensively for years to rapidly amplify targeted DNA sequence regions, certain shortcomings limit its application in diagnostics and detection. For instance, PCR product analysis must be carried out after amplification, giving rise to an issue of post-amplification contamination and carry-over contamination (Heid et al., 1996). Most importantly, a high ratio of false positive results are often associated with PCR due to non-specific binding of the primers and the subsequent non-specific amplification of products. Recently a real-time PCR technology has emerged as a powerful diagnostic tool in both medical and agricultural fields.

Using real-time PCR, a fluorescent dye such as SYBR green can be incorporated into the reaction mixture and the fluorescent signals, generated from fluorescent dye binding to double stranded DNA products, can be detected directly by the optical module coupled with the thermocycler. The signals are processed by computer data analysis software for almost real-time calculation and on screen plotting. A new dimension of real-time PCR called Taqman assay further introduced a third oligonucleotide probe, labeled with 5' fluorescent reporter dye and 3' quenching dye, for signal detection (Livak et al., 1995; Basseler et al., 1995). In the Taqman system, the quenching dye on the 3' end captures the fluorescence from the 5' reporter dye so the intact probe itself does not produce strong signal. During amplification when the probe hybridized to complementary sequence within the amplified products, the 5'>3' exonuclease activity of the polymerase in the reaction system cleaves the probe, minimized the quenching effect and the emitted fluorescent signal from the 5' reporter dye can be detected by the optical module. An advantage of applying the Taqman system is that a double complementing sequence selection mechanism by both the primers and the probe is involved, therefore the false positive rate of the detection can be significantly cut down. So far, various Taqman real-time PCR-based detection approaches have been reported. However, reports on its application in the real food system are still limited. The greatest challenges are (i) effective extraction of DNA and RNA

from a system where microorganisms are mixed with the food matrix including bulk proteins, carbohydrates and fatty acids, (ii) selection of primer-and-probe sets that are specific for the target microorganisms and do not interaction with background microflora and food ingredients, and (iii) minimizing the influence of food ingredients and other chemical compounds in the food matrix on the action of enzymes involved in DNA extraction and amplification.

Our objective was to demonstrate the feasibility of the real-time PCR based detection technology for food industry applications. It is our understanding that due to the complication of various food systems, detection procedures likely need to be optimized for individual food commodities. In this study, we investigated the practicability of using the Taqman-based real-time PCR approach in detecting target microorganisms in juice products. Here we report the effectiveness of the Taqman-based detection system in rapid, specific and sensitive detection of spoilage A. acidocaldarius and A. acidoterrestris in juice products, using a primer-and-probe set specific for the shc gene encoding squalene-hopene cyclase.

Materials and Methods

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15 Bacterial strains and growth conditions.

The bacterial strains used in the study and their growth conditions were listed in Table VI. ATCC 573 medium consists of 1.3g (NH₄)₂SO₄, 0.37g KH₂PO₄, 0.25g MgSO₄·7H₂O, 0.07g CaCl₂·2H₂O, 1.0g glucose, 1.0g yeast extract, and 1.0 L distilled H₂O, pH 4.0. BAM-SM ATCC 1656 medium consists of 0.25g CaCl₂·2H₂O, 0.5g MgSO₄·7H₂O, 0.2g (NH₄)₂SO₄, 3.0g KH₂PO₄, 6.0g yeast extract, 5.0g glucose, 1.0mL trace elements (0.66g CaCl₂·2H₂O, 0.18g ZnSO₄·7H₂O, 0.16g CuSO₄·5H₂O, 0.15g MnSO₄·4H₂O, 0.18g CoCl₂·6H₂O, 0.10g H₃BO₃, 0.30g Na₂MoO₄·2H₂O, 1.0L distilled H₂O), and 1.0L distilled H₂O. *Geobacillus stearothermophilus* ATCC 10149 was grown in Nutrient broth (Difco). Stock cultures of all strains were stored in their respective media plus 40% glycerol and kept at -80°C. All inoculations used were 2% concentrations made from frozen cultures.

Table VII. Bacteria cultures used in the study.

	Strains	Medium and Growth Condition	Resource	_
	A.acidocaldarius ATCC43030	#573 broth ^a at 48°C	ATCC	_
30	A.acidoterrestris ATCC49025	#1655 broth ^a at 48°C	ATCC	
	A.cycloheptanicus ATCC49029	#1656 broth ^a at 48°C	ATCC	
	Bacillus subtilis Geobacillus?	Nutrient broth ^b , 40°C		
	E. coli DH5α	LB broth, Miller ^c at 37°C		
35	Pseudomonus putidis?	LB broth, Miller at 37°C		
	Listeria monocytogenes V7 Tryptic soy broth ^d at 37°C			
	Lactococcus lactis 2301	M17 broth ^e at 37°C		
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^aAll numbered broth for *Alicyclobacillus* spp. are ATCC media.

^bFrom Becton Dickison & Co., Sparks, MD.

^cFrom Fisher Chem., Fais Lawn, NJ.

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^dFrom Becton Dickison and Company, Sparks, MD.

^eFrom Becton Dickison and Company, Sparks, MD.

DNA extraction, gene cloning and DNA sequencing. For DNA extraction, cells were collected from 1 ml of bacterial culture by micro-centrifugation 7.6K rpm for 10 min. The cell pellet was treated with 20 mg/ml of lysozyme (Sigma Chemical CO. St Louis, MO 63178, USA) in buffer for 45 min at 37°C. Genomic DNA was extracted using the DNeasy ® Tissue Kit (QIAGEN GmbH, D-40734 Hilden, Germany) and eluted into 100 μl of elution buffer following the instructions from the manufacturer.

The *shc* gene fragment from each strain was obtained by conventional PCR amplification using degenerate primers derived from conserved amino acid sequences and the genomic DNA from each strain as template. The reaction mixture includes 1X PCR buffer, 3mM MgCl₂, 4mM dNTP (Invitrogen, Carlsbad, CA), 1µM primer pairs, 1µl of genomic DNA template and ddH₂O in a total final volume of 50µl. PCR was performed one cycle at 95°C for 3min, followed by 30 cycles at 95°C for 30s, 50°C for 30s and 72°C for 1min, with a final extension at 72°C for 7min using I-cycler (Bio-Rad, Hercules, CA). PCR products were purified using the QIAquick PCR purification kit (Qiagen, Valencia, CA) following manufacturer's instruction. Purified PCR products were cloned into pCR 2.1 vectors and transformed into One Shot competent *Escherichia coli* cells using the TA Cloning kit (Invitrogen, Carlsbad, CA). Recombinant plasmids were recovered using QIAGEN miniprep (QIAGEN GmbH, D-40734 Hilden, Germany). DNA sequences were determined using the ABI PRISM® 3700 DNA Analyzer (Applied Biosystems, Foster City, CA) at the Plant Genome Sequence Facility, The Ohio State University.

Real-time Taqman PCR conditions For real-time PCR, the reaction was conducted in thin-wall microcentrifuge tubes including $1X iQ^{TM}$ Supermix (Bio-Rad, Hercules, CA), $0.5 \mu M$ of primer pair, $0.3\mu M$ of probe, $10\mu l$ of genomic DNA extraction and ddH_2O in a final volume of $50\mu l$.

PCR was performed one cycle at 95°C for 3min followed by 40 cycles of 95°C for 30s, 55°C for 1min using I-cycler (Bio-Rad, Hercules, CA).

<u>DNA sequence analysis.</u> The DNASTAR (DNASTAR, Madison, WI) software package was used in DNA and protein sequence alignment and homology search. DNA oligonucleotide primer and probe sequences were also compared with sequences from the GenBank sequence database using BlastSearch.

Specificity and sensitivity analyses Assays were conducted to test the specificity of the detection system against spoilage Alicyclobacillus spp. and other common food-borne microorganisms. Genomic DNA was extracted from broth cultures of A. acidoterrestris and A. acidocaldarius, grown for 48 h at 48°C (absorbance at OD₆₀₀ around 0.5-0.7), using the previously discussed DNA extraction protocol. Genomic DNAs extracted from 1 ml of overnight culture of Escherichia coli DH-5α, Lactococcus lactis subsp. lactis C2, Geobacillus stearothermophilus ATCC 10149 and Pseudomonas putida 49L/51 were also used in the specificity study. Ten micro liters out of the 100 micro liter of elution was used as template and the real-time PCR amplification was carried out using conditions described above but using 32 instead of 40 cycles of amplification.

The sensitivity tests of the real-time PCR assay for detection of Alicyclobacillus in bacterial culture media were performed using tenfold serial dilutions from 10^0 to 10^{-8} of A. acidoterrestris in a 10ml solution of 0.85% NaCl. The initial cultures were obtained by grown for 18 h at 48°C using 2% inoculation from the frozen stock, with the absorbance reading at OD_{600} range between 0.38 and 0.42. After serial dilution, cells from 1ml of each sample were collected by centrifugation at 7600 rpm for 10 minutes for DNA extraction. Ten microliter out of the 100 microliter of elution was used as template and the real-time PCR amplification was carried out as described above.

Sensitivity tests in juice products were also performed in the same manner but the serial dilutions were carried in apple juice instead of saline.

In both sensitivity analyses, a range of dilutions between 10⁻⁴ and 10⁻⁵ were plated on acidified PDA agar (Difco, Detroit) for colony counting to compare with the results by Taqman real-time PCR. Plates were incubated at 48°C for 48h.

Results

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25 1. The primer-and-probe set used in the real-time PCR Tagman assay.

Hopanoids are membrane components involved in maintaining membrane fluidity and stability (4) of *Alicyclobacillus* spp. in extreme environmental conditions. We have targeted the *shc* gene encoding squalene-hopene cyclase, a key enzyme in hopanoid biosynthesis, for PCR primer-and-probe development.

Using an established approach (Wang et al., 2001), squalene-hopene cyclase protein sequences from several microorganisms.were aligned and conserved amino acid sequences in squalene-hopene cyclase were identified. Figures 5 and 6, respectively, show the polynucleotide and protein alignments for two strains of Alicyclobacillus. Two degenerate primers 5' GGNGGNTGGATGTTYCARGC 3' (Y=C+T; R=A+G; N=A+T+C+G) (SEQ ID NO 64) and 5'

YTCNCCCCANCCNCCRTC 3' (SEQ ID NO 65) were derived. Using this set of primers and the genomic DNA from A. acidocaldarius ATCC 43030 and A. acidoterrestris ATCC 49025, the 705 bp shc fragments were amplified by PCR from both strains. The PCR fragments were cloned into the TA vector and the inserted DNA sequences were determined. The DNA sequences were further compared with other Alicyclobacillus spp. shc sequences in the GenBank. Three conserved oligonucleotides were derived including the Forward Primer ATGCAGAGYTCGAACG 3' (SEQ ID NO 25) and the Reverse Primer AAGCTGCCGAARCACTC 3' (SEQ ID NO 27) flanking a 136 bp fragment, and the Probe 5'TCRGARGACGTCACCGC3' (SEQ ID NO 26). The synthesized primers were ordered from Sigma-Genosys (The Woodlands, TX). The Probe is fluorescence-labeled with 5' 6-FAM BHQ-13' by Biosearch Technologies, Inc. (Novato, CA) and was used in the Taqman assay.

Specific detection of spoilage Alicyclobacillus spp.

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Real-time PCR assays were performed to determine the specificity of the primers and probe for spoilage Alicyclobacillus spp. E. coli DH-5a, L. lactis subsp. lactis C2, and P. putida 49L/51, G. stearothermophilus ATCC 10149 were also included in the study to test the possibility of crossreactions by the primer-and-probe set with common food-borne microorganisms. Assays were performed in triplicate, and a representative real-time PCR curve plotted by the iCycler iQ Optical System Software is shown in Figure 26.

Representative strains from A. acidocaldarius and A. acidoterrestris were tested positive. No cross-reaction was detected in other commonly found food-borne microorganisms. Further specificity study was conducted by searching the Blast databases for DNA sequences from the National Center for Biotechnology Information (NCBI). We found no combination of the above three oligonucleotides in other microorganisms but A. acidocaldarius and A. acidoterrestris. The data suggested that the system is specific for spoilage A. acidocaldarius and A. acidoterrestris.

25 Levels of detection in bacterial culture medium and in apple juice.

To establish the detection level using the above real-time PCR system, we have conducted 10⁰ to 10⁻⁶ serial dilutions of A. acidoterrestris ATCC 49025 in culture medium. Cells from 1 ml of diluted samples were collected and 10/100 of the DNAs extracted were used as template in the real-time PCR analysis. All experiments were repeated for at least three times and a representative curve was presented as Figure 27. Our results showed that using the above primer-and-probe set, the presence of as few as 10 cells cells in a sample could be detected. This detection level is comparable to results from other microbial detection studies using real-time PCR.

To further verify the feasibility of using the detection system in juice products, we have conducted 10^0 to 10^{-6} serial dilutions of A. acidoterrestris ATCC 49025 in apple juice. The experiments were repeated for three times and a representative curve was presented as Figrue 28. Similar detection level was achieved in apple juice.

2.Discussion and Conclusion

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Rapid, specific and sensitive detection of microorganisms in agricultural and food systems has proved to be a challenge. There are several major hurdles for effective microbial detection in the food systems. First, problematic food is normally associated with low level of initial contamination. However, the rich food matrix can support the growth of microbial agents in many cases during food storage and distribution. Thus even low level of initial contamination can cause serious damage. To be able to detect the presence of this low level contamination from food matrix often involving bulk proteins, carbohydrates and fatty acids, proper sampling and lengthy pre-detection enrichment steps are often required. To achieve rapid detection, predetection enrichment procedures need to be minimized and the detection system also should be sensitive enough to pick up low level of contamination.

Second, both foods and farm environment are complex ecosystems with significant background microflora. In addition to the background microflora normally associated with raw materials, beneficial microorganisms such as starter cultures sometimes are intentionally inoculated and present in large quantity in certain products. Therefore, to avoid false positive results, detection method for spoilage or pathogenic organisms needs to be specific enough to pick up only the target microorganisms. Finally, the rich and complex food ingredients often include various salts, carbohydrates, preservatives, emulsifiers, fatty acids, and proteins. The presence of these components varies among food commodities and can interfere with detection in various degrees. Therefore detection approaches and procedures need to be verified for effectiveness in these food systems.

Real-time Taqman PCR-based approach has the potential to achieve rapid, sensitive and specific detection. An average DNA amplification cycle for a small fragment can be completed within a minute. Theoretically after 30-40 cycles the amplification products from one DNA template in the system can be readily detected and plotted on the screen in almost real-time. The double sequence selection mechanism involving both the oligonucleotide primers and probe further minimizes the possibility of false positive results and enhances the detection specificity.

In this study, using a primer-and-probe set targeting the spoilage A. acidocaldarius and A. acidoterrestris, we were able to achieve specific detection without cross-reacting with representative strains from other common food-borne microorganisms including a strain from the

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closely related thermophilic G. stearothermophilus. Although only a few representative strains were used in the laboratory specificity studies, a computer-based search covering all the world-wide deposited DNA sequences available through the NCBI website was conducted to ensure that the combination of the sequences of the oligonucleotide primers and probe used in the study are distinctive enough to detect only A. acidocaldarius and A. acidoterrestris strains.

The level of detection limit with confidence is important for any detection approaches. In this study we have conducted sensitivity tests in both bacterial cultural medium and a real food system-apple juice. For laboratory handling purpose and for the convenient of using commercially available yet economically feasible DNA extraction kit, bacterial cells were serially diluted in either medium or juice and cells in 1 ml of samples were collected by microcentrifugation. DNA were extracted and 10/100 of the elution were used as template in PCR. The experiment was repeated at least three times and a representative curve presented as Figure 27. The lowest detection limit was determined based on the cell count numbers from agar plates derived from dilution with the optimal counting numbers (30-300) and the fold of dilution corresponding to each positive curves presented. Using this approach, we report that the presence of as few as 10 cells per sample with confidence. Because during each independent repeats the 10-fold serial dilutions were conducted without knowing exactly how many cells were in 1 ml of samples, the standard deviation reflects this fact. To further narrow down the range of standard deviation of detection, serial dilutions within the range of 2-10 can be conducted so a more precise confident level can be possibly established. We did not extrapolate the results using In other referred paper sometimes a standard curve was established first for sensitivity analysis. Furthermore, in a quality control laboratory, a regular sample size is normally 25 ml instead of 1 ml. Theoretically, sample detection limits can further be improved as long as cells from 25 ml or even 100 ml of samples can be properly collected and re-suspended in 1 ml of solution to conduct DNA extraction.

We are in the process of establishing a rapid detection system for food industry applications (the CleanPlant system) and the real-time Taqman PCR is one of our preferred platforms. In order to apply this detection platform in juice related products, we need to establish the feasibility of using the system for raw material screening and final product monitoring. We have conducted the sensitivity test by spiking the Alicyclobacillus in apple juice purchased from local grocery stores and similar level of detection was achieved indicating the applicability of such a system in final product screening. Further, we have used this system to detect the presence of Alicyclobacillus in apple juice concentrates, which are considered raw materials for the processing facilities. Similar level of detection was achieved except diluting and rinsing

procedures need to be incorporated to minimize inhibitory effects by the concentrated food ingredients (data not shown). These data suggested that

Because the system we developed is based on recognition of the signature DNA sequence of microorganisms, it has high specificity and does not cross react with other food-borne microorganisms (Figure 26). The detection limit was achieved in both bacterial culture medium and apple juice. Since no inhibition to the reaction system was detected using samples collected from apple juice, we expect the sensitivity of the detection system can be further improved by including a pre-treatment procedure to apply a centrifugation or membrane filtration procedure to concentrate the bacteria cells from a large sample volume. This approach is in fact a preferred practice in the industry where the sampling size varies from 25 ml to 1 liter. Since only 1/10 of the DNA extract was used in the reaction, we expect further improvement for the sensitivity can be achieved by incorporating more DNA template to the reaction system.

Example 3:

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Yeast genomic DNA extraction protocol:

Innoculate yeast, overnight; Centrifuge 10,000 rpm for 10 mins; Discard supernatant, add 600 ul Sorbital buffer (1 M Sorbital, 100 mM EDTA, 14 mM B-mercaptoethanol, 30 ul 20 mg/ml lyticase) in pellet, vortex, room temperature for 30 min; Centirfuge 10,000 rpm for 5 min; Add 180 ATL (Qiagen DNAeasy kit) and 20 ul proteinase K (Qiagen DNAeasy kit) to pellet and vortex; 55° for 1h, add 200 ul AL (Qiagen DNAeasy kit), 70° for 10 min; 200 ul Ethanol, vortex, apply to DNeasy spin column.; centrifuge 10,000 rpm for 1 min, discard flow-through' add 500 ul Buffer AW1 (Qiagen DNAeasy kit), spin for 1 min; add 500 ul Buffer AW1 (Qiagen DNAeasy kit), spin for 1 min.

Mold genomic DNA extraction protocol:

Innoculate Mold in PDB; 3 days later, centrifuge 10,000 rpm for 10 min; add 500 ul Mold extraction buffer (1% CTAB, 1.4 M NaCl, 100 mM Tris, 20 mM EDTA, pH 8.0) to pellet; 100 ul glass beads, water bath sonic (55°.) for 45 min; add 50 ul Proteinase K (Qiagen DNAeasy kit) and incubate in 55° for 1 h; Centrifuge 10,000 rpm for 5 min; Transfer the supernatant, add 500 ul AL (Qiagen DNAeasy kit), 70° for 10 min; Add 200 ul Ethanol and pipet it into Dneasy mini column; 10,000 rpm for 1 min; Add 500 ul AW1 (Qiagen DNAeasy kit), spin for 1 min; Add 500 ul AW2 (Qiagen DNAeasy kit), spin for 3 min; Add 100 AE buffer (Qiagen DNAeasy kit), spin for 1 min.

What is claimed is:

- 1. A method for detecting Alicyclobacillus and Geobacillus in a test sample, the method comprising
 - (a) providing an oligonucleotide set comprising:
- (i) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1;
- (ii) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1, said second consecutive sequence being downstream from said first consecutive sequence,; and
- (ii) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences;
- (b) amplifying DNA in the sample with the said primer set and a polymerase chain reaction, and
- (c) determining the presence of PCR products of step (b), wherein the presence of a PCR product in the sample is indicative of contamination of the test sample by *Alicyclobacillus* or *Geobacillus* or both.
- 2. The method of claim 1 wherein the forward primer has a sequence selected from the group consisting of SEQ ID NO 1, SEQ ID NO 5, SEQ ID NO 9, SEQ ID NO 13, SEQ ID NO 17, and SEQ ID NO 20.
- 3. The method of claim 1 wherein the reverse primer has a sequence selected from the group consisting of SEQ ID NO 4, SEQ ID NO 8, SEQ ID NO 12, SEQ ID NO 16, SEQ ID NO 19, and SEQ ID NO 23.
- 4. The method of claim 1 wherein the probe has a sequence selected from the group consisting of SEQ ID NO 2, SEQ ID NO 3, SEQ ID NO 6, SEQ ID NO 7, SEQ ID NO 10, SEQ ID NO 11, SEQ ID NO 14, SEQ ID NO 15, SEQ ID NO 18, SEQ ID NO 21, and SEQ ID NO 22.

5. A method for detecting Alicyclobacillus and Geobacillus in a test sample, the method comprising

- (a) providing a oligonucleotide set comprising:
- (i) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5;
- (ii) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5, said second consecutive sequence being downstream from said first consecutive sequence,; and
- (ii) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences;
- (b) amplifying DNA in the sample with the said primer set and a polymerase chain reaction, and
- (c) determining the presence of PCR products of step (b), wherein the presence of a PCR product in the sample is indicative of contamination of the test sample by *Alicyclobacillus* or *Geobacillus* or both.
- 6. The method of claim 5 wherein the forward primer has a sequence selected from the group consisting of SEQ ID NO 28, and SEQ ID NO 29.
- 7. The method of claim 5 wherein the reverse primer has a sequence selected from the group consisting of SEQ ID NO 34, SEQ ID NO 35, SEQ ID NO 36, and SEQ ID NO 37.
- 8. The method of claim 5 wherein the probe has a sequence selected from the group consisting of SEQ ID NO 30, SEQ ID NO 31, SEQ ID NO 32, and SEQ ID NO 33.
- 9. A method for detecting Alicyclobacillus and Geobacillus in a test sample, the method comprising
 - (a) providing a oligonucleotide set comprising:

(i) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 1;

- (ii) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 1, said second consecutive sequence being downstream from said first consecutive sequence,; and
- (ii) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 1, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences;
- (b) amplifying DNA in the sample with the said primer set and a polymerase chain reaction, and
- (c) determining the presence of PCR products of step (b), wherein the presence of a PCR product in the sample is indicative of contamination of the test sample by *Alicyclobacillus* or *Geobacillus* or both.
- 10. The method of claim 9 wherein the forward primer has a sequence selected from the group consisting of SEQ ID NO 38, SEQ ID NO 39, SEQ ID NO 40, SEQ ID NO 41, SEQ ID NO 42, and SEQ ID NO 43.
- 11. The method of claim 9 wherein the reverse primer has a sequence selected from the group consisting of SEQ ID NO 50, SEQ ID NO 51, SEQ ID NO 52, and SEQ ID NO 53.
- 12. The method of claim 9 wherein the probe has a sequence selected from the group consisting of SEQ ID NO 44, SEQ ID NO 45, SEQ ID NO 46, SEQ ID NO 47, SEQ ID NO 48, and SEQ ID NO 49.
- 13. A method for detecting mold or yeast in a test sample, the method comprising
 - (a) providing a oligonucleotide set comprising:
- (i) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8;

(ii) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8, said second consecutive sequence being downstream from said first consecutive sequence,; and

- (ii) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences;
- (b) amplifying DNA in the sample with the said primer set and a polymerase chain reaction, and
- (c) determining the presence of PCR products of step (b), wherein the presence of a PCR product in the sample is indicative of contamination of the test sample by *mold or yeast* or both.
- 14. The method of claim 13 wherein the forward primer has a sequence selected from the group consisting of SEQ ID NO 54, and SEQ ID NO 58.
- 15. The method of claim 13 wherein the reverse primer has a sequence selected from the group consisting of SEQ ID NO 55, and SEQ ID NO 59.
- 16. The method of claim 13 wherein the probe has a sequence selected from the group consisting of SEQ ID NO 56, SEQ ID NO 57, and SEQ ID NO 60
- 17. A method for detecting mold or yeast in a test sample, the method comprising
 - (a) providing a oligonucleotide set comprising:
- (i) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8;
- (ii) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8, said second consecutive sequence being downstream from said first consecutive sequence,; and

(ii) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences;

- (b) amplifying DNA in the sample with the said primer set and a polymerase chain reaction, and
- (c) determining the presence of PCR products of step (b), wherein the presence of a PCR product in the sample is indicative of contamination of the test sample by *mold or yeast* or both.
- 18. The method of claim 17 wherein the forward primer has a sequence of SEQ ID NO 61.
- 19. The method of claim 17 wherein the reverse primer has a sequence of SEQ ID NO 62.
- 20. The method of claim 17 wherein the probe has a sequence of SEQ ID NO 63.
- 21. The method of claim 1 wherein the primers
 - i.) do not contain runs of more than 5 of the same nucleotide base,
 - ii) do not contain internal palindromic sequences,
 - iii) do not hybridize to one another under stringent conditions, and
 - iv) have 40 to 60 percent G+C content, and

wherein said PCR amplification provides a PCR product that is from 50 to 613 nucleotides in length

- 22. The method of claim 1, wherein the PCR is quantitative PCR.
- 23. The method of claim 1, wherein the PCR is real-time PCR.
- 24. A method of detecting the presence of acidic bacteria in a test sample using real time monitoring of a polymerase chain reaction amplification of a target nucleic acid sequence found in the acidic bacteria, said method comprising the steps of
- (a) adding to the test sample an effective amount of a forward nucleic acid primer and reverse nucleic acid primer and a nucleic acid probe, wherein the forward primer is selected from the group consisting of SEQ ID NO 1, SEQ ID NO 5, SEQ ID NO 9, SEQ ID NO 13, SEQ ID NO 17, and SEQ ID NO 20, and wherein the reverse primer is selected from the group consisting of SEQ ID NO 4, SEQ ID NO 8, SEQ ID NO 12, SEQ ID NO 16, SEQ ID NO 19, and SEQ ID NO 23, and wherein the probe is selected from the group consisting of SEQ ID NO 2, SEQ ID NO 3,

SEQ ID NO 6, SEQ ID NO 7, SEQ ID NO 10, SEQ ID NO 11, SEQ ID NO 14, SEQ ID NO 15, SEQ ID NO 18, SEQ ID NO 21, and SEQ ID NO 22 and wherein the probe hybridizes to an amplified copy of the target nucleic acid sequence, and wherein the probe is labeled with a marker which emits a signal upon the hybridization of the probe to the target nucleic acid sequence;

- (b) amplifying the target nucleic acid sequence by polymerase chain reaction;
- (c) detecting the emitted signal of the sample.
- 25. A method of detecting the presence of fungi in a test sample using real time monitoring of a polymerase chain reaction amplification of a target nucleic acid sequence found in the acidic bacteria, said method comprising the steps of
- (a) adding to the test sample an effective amount of a forward nucleic acid primer and reverse nucleic acid primer and a nucleic acid probe, wherein the forward primer is selected from the group consisting of SEQ ID NO 54, and SEQ ID NO 58 and SEQ ID NO 61
- , and wherein the reverse primer is selected from the group consisting of SEQ ID NO 55, SEQ ID NO 59, and SEQ ID NO 62, and wherein the probe is selected from the group consisting of SEQ ID NO 56, SEQ ID NO 57, SEQ ID NO 60, and SEQ ID NO 63 and wherein the probe hybridizes to an amplified copy of the target nucleic acid sequence, and wherein the probe is labeled with a marker which emits a signal upon the hybridization of the probe to the target nucleic acid sequence;
- (b) amplifying the target nucleic acid sequence by polymerase chain reaction;
- (c) detecting the emitted signal of the sample.
- 26. A kit for detecting *Alicyclobacillus* and *Geobacillus* in a test sample, comprising a set of oligonucleotides comprising:
- (a) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1;
- (b) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1, said second consecutive sequence being downstream from said first consecutive sequence; and

(c) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 1327 through nucleotide position 1460 shown in Figure 1, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences.

- 27. A kit for detecting *Alicyclobacillus* and *Geobacillus* in a test sample, comprising a set of oligonucleotides comprising:
- (a) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5;
- (b) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5, said second consecutive sequence being downstream from said first consecutive sequence; and
- (c) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 334 through nucleotide position 485 shown in Figure 5, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences.
- 27. A kit for detecting *Alicyclobacillus* and *Geobacillus* in a test sample, comprising a set of oligonucleotides comprising:
- (a) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 5;
- (b) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 5, said second consecutive sequence being downstream from said first consecutive sequence; and
- (c) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 752 through nucleotide position 813 shown in Figure 5, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences.

28. A kit for detecting *yeast or mold* in a test sample, comprising a set of oligonucleotides comprising:

- (a) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8;
- (b) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8, said second consecutive sequence being downstream from said first consecutive sequence; and
- (c) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 81 through nucleotide position 225 shown in Figure 8, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences.
- 29. A kit for detecting *yeast or mold* in a test sample, comprising a set of oligonucleotides comprising:
- (a) a forward primer of from 15 to 35 nucleotides in length, said forward primer comprising a sequence which is identical to a first consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8;
- (b) a reverse primer of from 15 to 35 nucleotides in length, said reverse primer comprising a sequence which is complementary to the inverse complement of a second consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8, said second consecutive sequence being downstream from said first consecutive sequence; and
- (c) a probe of from at least 15 nucleotides in length, said probe comprising a sequence which is which is identical to a third consecutive sequence within the sequence of nucleotide position 114 through nucleotide position 238 shown in Figure 8, or the complement thereof, and falls between or overlaps with the first and second consecutive sequences.

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560 538 557	AAGGGTGCGTAGGCGGTCGAGCAAGTCTGGAGTGAAGTC 43030 168 AAGCGTGCGTAMGCGGTTGTGTAAGTCTGAAAGTC 49025 168 AAGCGTGCGTAGGCGGTTGCGTGTGTCCGGGGTGAAAGTC 49029 168	
	CA. GGCTC. ACC. TGGG GC. TTGGAAACTGC. T AC Consensus #1 CAXGGCTCAACCXTGGGAATGCTTTGGAAACTGCXTG - AC Majority	
600 578 597	SCTTTGGAAACTGCTTG-A GCTTTGGAAACTGCATGGA GCCTTGGAAACTGCGTA-A	

Δligni ∹rida	Alignment Report of Untitled ClustalV (Weighted) Friday, September 05, 2003 9:42 AM	Page
	TTGAGTGCTGGAGAGGC. AGG AATTCC. CGTGT. A. CG Consensus #1 TTGAGTGCTGGAGAGGGGGAATTCCXCGTGT - AGCG Majority	
	650 660 670 680	
639 618 636	TTGAGTGCTGGAGAGGCAAGGGGAATTCCACGTGT-AGCG 43030 16s TTGAGTGCTGGAGAGGCNAGGCNAATTCCNCGTGTTACCG 49025 16s TTGAGTGCTGGAGAGGCAAGGGGAATTCCGCGTGT-AGCG 49029 16s	
	CCAGTGGC.	
678 658 675	T - AGAGAT SNTANATAT T - AGATAT	
	A.GCGCCTT.GCTGGACAGTG.ACTGACGCTGA.GGCACG Consensus #1 AXGCGCCTT-GCTGGCAGTG-ACTGACGCTGA-GGCACG Majority 730 740 750 750	
716 698 713	ARGCGCCTT - GCTGGACAGTG - ACTGACGCTGA - GGCACG 43030 168 ANGCGCCTTTGCTGGACAGTGGACTGACGCTGAAGGCACG 49025 168 AGGCGCCTT - GCTGGACAGTG - ACTGACGCTGA - GGCACG 49029 16s	
	AAA-GCGTGGGGAGCAA	
753 738 750	GGAGCAA	

. Align Frida	Alignment Report of Untitled ClustalV (Weighted) Friday, September 05, 2003 9:42 AM		Page 6
		# J	
	810 820 830 840		
769	ACAGGATTAGATACCCTG - GTAGTCC - ACG 43030 1 AAACAGGATTAGATTCCCNTTGTAGTCCCGCC 49025 1	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	
90/	T SOCA T T W G A I A C C C I G T G G G G G G G G G G G G G G G G G		
	CGTAAACGATGAGTGCT-AGGTGTTGGGGGGACA	T ⊭	
	088 028 098 058		
797	CCGTAAACGATGAGTGCT-AGGTGTTGGGGGGGACACACCC 43030 16	16s 16s	
794	CGTAAACGATGAGTGT-AGGTGTTGGGGGGTACCACC 49029	89	
	CA. TGC. G GGAAA. CCAATAAGCACTCCGCCTGGGGA Consensus	T# 8:	
	- CAGTGCCG A A G G A A A C C C A A T A A G C A C T C C G C C T G G G G A Majority		
	890 900 910 920	•	
836	- CAGTGCCGAAGGAAAMCCAATAAGCACTCCGCCTGGGGA 43030 1	68	
858	CANTGC - GGNGGAAACCCAATAAGCACTCCGCCTGGGGA 49025		
833	TCAGTGCGGAAGGAATAAGCACTCCGCTGGGGA 49029 1	8 9	
	•	18 #1	
	GTACGGTCGCAAGACTGAAACTCAAAGGAATTGACGGGGG Majority		
	930 940 950 960	·.	
875	GTACGGTCGCAAGACTGAAACTCAAAGGAATTGACGGGGGG43030 1	. 89	
873	TACGGTCGCAAGACTCAAAGGAATTGACGGGG		

Align Frida	Alignment Report of Untitled ClustalV (Weighted) Friday, September 05, 2003 9:42 AM	Page 7
	CCCGCACAAGCAGTGGAGCAT. TGGTTTAA. TCGAAGCAA Consensus #1	
	0001 066 086 026	
915 936	CCCGCACAAGCAGTGGAGCATGTGGTTTAAATCGAAGCAA 43030 168	
913	CCCGCACAAGCAGCATGTGGTTTAATTCGAAGCAA 49029 16s	·
	CGCGAAGAACCTTA. CAGGGCT. GACATCCC. CTGAC Consensus #1	
	1010 1020 1030 1040	•
955 976 953	CGCGAAGAACTTACCAGGGCTTGACATCCCTTGACACGGGGTCGCCTCTGACACGGGGCTNGACATCCCTCTGACCGGG 49025 168	
	CGCAGAGATGTCCCTTCGGGGCAG.GGAGACAGGT Consensus #1	
	1050 1060 1070 1080	
995	CTCAGAGATGAGGGTCCCTTCGGGGCAGAGACAGT 43030	
933 933	CGCAGAGATGCGGTTTCCCTTCGGGGCAGGGACAGGT	
	GGTGCATGGTTGTCAGCTCGTGTGAGATGTTGGG Consensus #1 GGTGCATGGTTGTCGTCAGCTCGTGTGAGATGTTGGG Majority	
	1120	
1035 1056 1033	GGTGCATGGTTGTCGTCAGCTCGTGTGTGAGATGTTGGG 43030 16s GGTGCTTGCATGTTGGG 49025 16s GGTGCATGTTGGG 49029 16s	

Alignment Report of Untitled ClustalV (Weighted) Alignment 1 Friday, September 05, 2003 9:42 AM	Page 8
TT. AGTCCCGCAACGAGCGCAACCCTTGA. CTGTGTTACC Consensus #1 TTAAGTCCCGCAACGAGCGCAACCCTTGAXCTGTGTTACC Majority	
1130 1140 1150 1160	
1075 TTCAGTCCCGCAACGAGCGCAACCCTTGACCTTGTGTTACC 43030 16s 1096 TTAAGTCCCGCAACGCAACCCTTGATGTTACC 49025 168	
1073 TTAAGTCCCGCAACGAGCGCAACCCTTGAACTGTGTTACC 49029 168	
AGC. CGT GG. GGGGACTCACAG. TGACTGCCGGCGTA Consensus #1 AGCACGTTGAGGTGGGGACTCACAGGTGACTGCCGGCGTA Majority	
1170 1180 1190 1200	
1115 A G C G C G T T G A G G C G G G G A C T C A C A G G T G A C T G C C G G C G T A 43030 165 1136 A G C A C G T T G T G G G G A C T C A C A G G T G A C T G C C G G C G T A 49025 165 1113 A G C A C G T G A A G G T G G G G A C T C A C A G T T G A C T G C C G G C G T A 49029 16s	
AGTCGGAGGAAGGCGGGATGACGTCAAATCATGACCC Consensus #1	
1210 1220 1230 1240	
1155 AGTCGGAGGAAGGCGGGATGACGTCAAATCATGCCC 43030 168 1176 AGTCGGAGGAGGGATGACGTCAAATCATGCCC 49025 168	
1153 AGTCGGAGGAAGGCGGGATGACGTCAAATCATGCCC 49029 16s	
TITATGICCIGGCTACACGTGCTACAATGGGCGG.AC Consensus #1	
1250 1260 1270 1280	
1195 CTGATGTCCTGGGCTACACGTGCTACAATGGGCGGAAC 43030 16s 1216 TTTATGTCCTGGGCTACACGTGCTACAATGGGCGGTAC 49025 16s 1193 TTTATGTCCTGGGCTACACGTGCTACAATGGGCGGTAC 49029 16s	

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Alignment Report of Untitled ClustalV (Weighted) Friday, September 05, 2003 9:42 AM	Page 9
AA.GGGA.GCGACCGCGAGG.GGAGC.AA.CCAAA Consensus #1	
1290 1300 1310 1320	
AAAGGGAAGGCGAAGCCGAGGCGGAAACCCAAAA 43030 AACGGGAAGCCGCGAGGTGGAGCAAAACCTAAAA 49025	
CG.TCGTAGTTCGGATTGCAGGCTGCAACTCGCTGC	
1340 , 1350 1360	
1275 G C C G C T C G T A G T T C G G A T T G C A G G C T G C A A C T C G C C T G C A 43030 16s 1273 G C C G T T C G T A G T T C G G A T T G C A G G C T G C A A C T C G C T G C A 49029 16s	
AATTGCTAGTAATCGCGGATCAGCATGCCG	
1315 TGAAGCCGGAATTGCTAGTAATCGCGGATCAGCATGCCGC 43030 16s 1336 TGAAGCCGGAATTGCTAGTAATCGCGGATCAGCATGCCGC 49025 16s 1313 TGAAGCCGGAATTGCTAGTAATCGCGGATCAGCATGCCGC 49029 16s	•
GGTGAAT.CGTTCCCGGGCCTTGTACACACGCCCGTCAC Consensus #1 GGTGAATCCGGGGCCTTGTACACACGCCCGTCAC Majority 1410	
TCCCGGGCCTTGTACACACGGCCCGTCA	

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(Weighted) Alignment I C G C A A C C C G A A G T C G G T G A G G T A A C C Consens C G G C A A C C C G A A G T C G G T G A G G T A A C C Majorit 14f0 (4 t ⁰) 1470 1480 C G C A A C A C C G A A G T C G G T G A G G T A A C C 49025 1 C G G C A A G T C G G T G G G T A A C C 49025 1 C C A G C C G A A G G T G G G T G G G T A A C C 49025 1 C C A G C C G A A G G T G G G G T G A T G A T T G A T T G A T G A T T G A T T G A T T G A T T G A T T G A T G A T		Ω.	a a a	Ľ2,	22 23 22 22 23 23	m,	8 8 8 9 9
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Consensus 'Consensus #1': When all match the residue of the Consensus show the the Consensus, otherwise show residue of

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49029

AGAGTTTGATCCTGGCTCAGGACGACGCTGGCGGCGTGCCTAATACATGCAAGTCGAGCGGACCCTTCGGGGTCAGCGG CGGACGGGTGAGTAACACGTGGGTAATCTGCCCAACTGACCGGAATAACGCCTGGAAACGGGTGCTAATGCCGGATAGGC **3GGTAACGGCTCACCAAGGCGACGATGCGTAGCCGACCTGAGAGGGTGGACGGCCACACTGGGACTGAGACACGGCCCAG** CCTTCGGGTTGTAAAGCTCAGTCACTCGGGAAGAGCGGCAAGGGGGAGTGGAAAGCCCCTTGAGAGACGGTACCGAGAGA CTGGGCTACACACGTGCTACAATGGGCGGTACAACGGGAAGCGAGACGGGGAGGTGGAGCCAAACCCCTGAAAGCCGTTCG ACTCCTACGGGAGGCAGCAGTAGGGAATCTTCCGCAATGGGCGCAAGCCTGACGGAGCAACGCCGCGTGAGCAAGAAGA SAAGCCCCGGCTAACTACGTGCCAGCAGCCGCGGTAATACGTAGGGGGCAAGCGTTGTCCGGAATCACTGGGCGTAAAGC STGCGTAGGCGGTTGCGTGTGTCCGGGGTGAAAGTCCAGGGCTCAACCCTGGGAATGCCTTGGAAACTGCGTAACTTGAG IGCTGGAGAGGCCAAGGGGAATTCCGCGTGTAGCGGTGGAATGCGTAGATATGCGGAGGAATACCAGTGGCGAAGGCGCCT rgctggacagtgactgacgctgaggcacgaaagcgtggggagcaaacaggattagataccctggtagtccacgccgtaaa CGATGAGTGCTAGGTGTTGGGGGGTACCACCCTCAGTGCCGAAGGAAACCCAATAAGCACTCCGCCTGGGGGAGTACGGTC GCAAGACTGAAACTCAAAGGAATTGACGGGGCCCCGCACAAGCAGTGGAGCATGTGGTTTAATTCGAAGCAACGCGAAGA ACCITIANCAGGGCTCGACATCCCCCTGACAGCCGCAGAGATGCGGTTTCCCTTCGGGGGAGGGGAGACAGGTGGTGCATG GTTGTCGTCAGCTCGTGTCGTGAGATGTTGGGTTAAGTCCCGCAACGAGCGCAACCCTTGAACTGTGTTTACCAGCACGTG **TAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGCCGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATC** CGTTCCCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGTCGGCAACACCCGGAAGTCGGTGGGGTAACCCGTCAGGG **AAGGTGGGGACTCACAGTTGACTGCCGGCGTAAGTCGGAGGAAGGCGGGGATGACGTCAAATCATGACGTTATGT**C **AGCCAGCCGCCGAAGGTGGGGTTGATGATTGGGGTGAAGTCGTAACAAGGTAGCCGT** Sequence, S 15 20 10

Figure 3

	4 V U S J L U	
	GACGAACGCTGGCGGCGTGCCTAATACATGCAAGTCGAGCGAG	80
2	GGGCAATCCGCCTTTCAGACTGGAATAACACTCGGAAACGGGTGCTAATGCCGGATAATACACGGGTAGGCATCTACTTG 1	160
	TGTTGAAAGATGCAACTGCATCGCTGAGAGAGGAGCCCGCGCGCG	240
	GACGATGCGTAGCCGACCTGAGAGGGTGACCGGCCACACTGGGACTGAGACACGCCCAGACTCCTACGGGAGGCAGCAG	320
	TAGGGAATCTTCCGCAATGGGCGCAAGCCTGACGGAGCAACGCCGCGTGAGCGAGAAGAAGGCCTTCGGGTTGTAAAGCTCT 4	400
	GTTGCTCGGGGAGAGACGACAAGGAGAGTGGAAAGCTCCTTGTGAGACGGTACCGAGTGAGGAGGCCCCGGCTAACTACGT 4	480
01	GCCAGCAGCCGCGGTAATACGTAGGGGGCAAGCGTTGTCCGGAATCACTGGGGCGTAAAGCGTGCGT	260
	AGTCTGAACTGAAAGTCCAAGGCTCNACCTTGGGNATGCTTTGGAAACTGCATGGACTTGAGTGCTGGAGAGGCNAGGCN	640
	AATICCNCGIGITACCGGIGNAAAIGCGNTANATAIGIGGAGGAATACCAGIGGCNAANGCGCCTIIGCIGGACAGIGGA 7	720
	CTGACGCTGAAGGCACGAAAANCGTGGGGANCAACNGGATTANATCCCCNAANGCGNGGGGAAGCAAACAGGATTAGATT 8	800
	CCCNTTGTAGTCCCGCCCCGTAANCNATGAGTACTTAGTTGTTGGGGGAACACACCCCANTGCGGNGGAAACCCAATAAG 8	880
15	CACTCCGCCTGGGGAGTGCGGTCNCAAGACTGAANCTCAAAGGAATTGACGGGGGCCCGCACAAGCAGTGGAGCATNTGG 9	096
	TITAATICGAAGCAACGCGAAGAACCITACCAGGGCINGACATCCCICTGACCGGTGCAGAGATGTACCTICCCTTCGGG 1	1040
	GCAGAGGAGACAGGTGGTGCATGGTTGTCGTCAGCTCGTGTGAGATGTTGGGTTAAGTCCCGCAACGAGCGCAACCC	1120
	TTGATCTGTGTTACCAGCACGTTGTGGTGGGGACTCACAGGTGACTGCCGGCGTAAGTCGGAGGAAGGCGGGGATGACGT 1	1200
	CAAATCATCATGCCCTTTATGTCCTGGGCTACACACGTGCTACAATGGGCGGTACAACGGGAAGCGAAGCCGCGAGGTGG 1	1280
20	AGCAAAACCTAAAAAGCCGTTCGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGCCGGAATTGCTAGTAATCGC 1	1360
	GGATCAGCATGCCGCGGTGAATCCGTTCCCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGTCGGCAACACCCGAA	1440
	GICGGIGAGGIAACCGIIAIGGAGCCAGCCGCCGAAGGIGGGGITGAIGAITGGGGIGAAGICGIAACAAGGIAGCCGI	1519

Figure 4

43030

GGCGGACGGGTGAGGAACACGTGGGTAATCTGCCTTTCAGGCCGGAATAACGCCCGGAAACGGGCGCGTAAAGCCGGATAC GCGGGGTAACGGCCCACCAAGGCGACGATGCGTAGCCGACCTGAGAGGGTGACCGGCCACACTGGGACTGAGACACGGCC CAGACTCCTACGGGAGGCAGCAGTAGGGAATCTTCCGCAATGGGCGCAAGCCTGACGGAGCAACGCCGCGCGTGAGCGAAGA <u>AGGCCTTCGGGTTGTAAAGCTCTGTTGCTCGGGGAGAGCGGCATGGGGGATGGAAAGCCCCGTGCGAGACGGTACCGAAGT</u> GAGGAAGCCCCGGCTAACTACGTGCCAGCAGCCGCGGTAAAACGTAGGGGGCGAGCGTTGTCCGGAATCACTGGGCGTAA GAGTGCTGGAGAGGCAAGGGGAATTCCACGTGTAGCGGTGAAATGCGTAGAGATGTGGAGGAATACCAGTGGCGAARGCG CCTTGCTGGACAGTGACTGACGCTGAGGCACGAAAGCGTGGGGAGCAAACAGGATTAGATACCCTGGTAGTCCACGCCGT <u>AAACGATGAGTGCTAGGTGTTGGGGGGACACCCCCAGTGCCGAAGGAAAMCCAATAAGCACTCCGCCTGGGGAGTACGG</u> TCGCAAGACTGAAACTCAAAGGAATTGACGGGGGCCCCGCACAAGCAGTGGAGCATGTGGTTTAAATCGAAGCAACGCGAA IGGTTGTCGTCAGCTCGTGTCGTGAGATGTTGGGTTCAGTCCCGCAACGAGCGCAACCCTTGACCTGTGTTACCAGCGCG TTGAGGCGGGGACTCACAGGTGACTGCCGGCGTAAGTCGGAGGAAGGCGGGGATGACGTCAAATCATCATGCCCTGATG TCCTGGGCTACACACGTGCTACAATGGGCGGAACAAAGGGAGGCGAAGCCGCGAGGCGGAGCGAAACCCCAAAAGCCGCT CGTAGTICGGATIGCAGGCTGCAACTCGCCTGCATGAAGCCCGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAA TACGITICCCGGGCCITIGTACACACCGCCCGTCACACCACGAGAGTCGGCAACACCCCGAAGTCGGTGAGGTAACCCCTGTG <u> AGAGTTTGA TCCTGGCTCAGGACGAACGCTGGCGGCGTGCCTAATACATGCAAGTCGAGCGGGTCTCTTCGGAGGCCAGC</u> <u> AGGGTGCGTAGGCGGTCGAGCAAGTCTGGAGTGAAAGTCCATGGCTCAACCATGGGATGGCTTTGGAAACTGCTTGAAACTGCTTT</u> 3GGAGCCAGCCGCCGAAGGTGGGTCGATGATTGGGGTGAAGTCGTAACAAGGTAGCCGT 15 20 S 10

Figure 5: Shc polynucleotide sequence alignments

Cloned ac 43030: Cloned Alicyclobacillus acidocaldarius ATCC43030
Cloned at 43030: Cloned Alicyclobacillus acidoterrestris ATCC49025
Blast ac: sequence of A. acidocaldarius got from the blast database
Blast at: sequence of A. acidocaldarius got from the blast database

Primer and probe ranges were highlighted red

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10		_			_	_	_	_	_							_	_	_		_	_		
																					C -+-	Majority	
											10										20		
15	1										•										C +-	cloned ac 43030	shc
																						Blast ac shc	
																						cloned at 49025 Blast at shc	shc
20	-																						
20																					G -+-	Majority	
											30										40		
`.	21										•											cloned ac 43030	shc
25	21																					Blast ac shc	
	21 21																					cloned at 49025 Blast at shc	shc
	2.																						
30																					G -+-	Majority	
	•										50)									60		
	41																				-+-	cloned ac 43030	shc
•			_																	•		Blast ac shc	
35																						cloned at 49025	shc
	41	A	Т	A	G	T	G	G	C	T	Т	G	A	Ċ	C	G	T.	C.	T	T.	G	Blast at shc	
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45	61																					Blast at shc	
		T	С	С	G	G	С	С	G	A	т	С	Α	т	-	G	С	С	G	G	G	Majority	
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	81																					cloned ac 43030	shc
	81																					Blast ac shc	- 1
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55	31	J	Č	_		_	•		-		•	-	••	-		_	_		•	_	_		

Figure 5: Shc polynucleotide sequence alignments (continued)

TTGGTTAAGGC--GGGTGAG Majority

5				-+	
				110	120
		TTG	G T C A A G G T C A A G	G C G G	C T G A A cloned ac 43030 shc G C G A G Blast ac shc
10	100 100				G T G A G cloned at 49025 shc G T G A G Blast at shc
		יה כו כי י	י יי כי יי כ	сстссс	C A G A T Majority
				-+	+-
15				130 -+	. 140
	121	TGG	CTGTTG	GACCGG	C A G A T cloned ac 43030 shc
	118 118				C A G A T Blast ac shc C A A A T cloned at 49025 shc
20	118				C A A A T Blast at shc
				TGGCGA	CTGGG Majority
				150	160
25	141			-+	TTGGG cloned ac 43030 shc
	138	CAC	3	G G G C G A	C T G G G Blast ac shc
					C T G G A cloned at 49025 shc C T G G A Blast at shc
30	138	1 (1)	CAAGGA	IGGCGA	
				G C C G G A	AGGTG Majority
				170	180
35	161			-+	A C C T C cloned ac 43030 shc
	158	C G G '	TGAAGC	GCCCGA	A C C T C Blast ac shc
	158 °				A G G C G cloned at 49025 shc A G G C G Blast at shc
	130				
40				G G T T T G	GCGTT Majority
				190	200
	181	AAC		-+	GCGCT cloned ac 43030 shc
45	178	AAG	CCGGGC	GGGTTC	G C G T T Blast ac shc
					G C A T T cloned at 49025 shc G C A T T Blast at shc
	170				
50				. C T G C G T	GTACT Majority
				210	220
	201	CCA	GTTCGA	CAACGT	G T A C T cloned ac 43030 shc
	198	CCA	GTTCGA	CAACGT	GTACT Blast ac shc
55	198 198	TGA.	A T T C C A A T T C C A	. C T G C G A	AAACT cloned at 49025 shc AAACT Blast at shc

Figure 5: Shc polynucleotide sequence alignments (continued)

		CCCGGACGTGGACGATACG Majority	
5		230 240	
10	218	TCCGGACGTGGACGACACG cloned ac 43030 s CCCGGACGTGGACGACACG Blast ac shc CCCAGACGTCGACGATACG cloned at 49025 s CCCAGACGTCGACGATACG Blast at shc	
		CGGTGGTCGTCTTGGCGCT Majority	
15		250 260	
20		CCGTCGTCATCTGGGCGCT cloned ac 43030 s CCGTCGTGGTGTGGCCT Blast ac shc CGATGGTCTTGGCGCT cloned at 49025 s CGATGGTCGTCTTGGCGCT Blast at shc	
20		AATGGCCTTCGATTGCCGG Majority	
		270 280	
25	258	AACACGCTGCGACTCCCGG cloned ac 43030 s AACACCCTGCGCTTGCCGG Blast ac shc AATGGCATTCAATTGCCGG cloned at 49025 s AATGGCATTCAATTGCCGG Blast at shc	
30		TGAGGGGCGGCGTCGTGAC Majority	
		290 300	
35	278	CGAGCGCCGCAGGCGAGAC cloned ac 43030 s CGAGCGCCGCAGGCGAC Blast ac shc TGAAGGGAAGCGTCGTGAC cloned at 49025 s TGAAGGGAAGCGTCGTGAC Blast at shc	
40		CCTTGACGCGTGGCTTCCG Majority	
		310 320	
45		C C A T G A C G A A G G G A T T C C G cloned ac 43030 s C C A T G A C G A A G G G A T T C C G Blast ac shc C A T T G A C C C G T G G C T T C C G cloned at 49025 s C A T T G A C C C G T G G C T T C C G Blast at shc	
		TGGTTTGTCGGGATGCAGA Majority	
50		330 340	
55	318 318	TGGATTGTCGGCAGA Cloned ac 43030 S TGGATTGTCGGCAGA Blast ac shc TGGTTGCGCGAGACLONED ATGCAGA Cloned at 49025 S TGGTTGCGCGAGACATGCAGA Blast at shc	

Figure 5: Shc polynucleotide sequence alignments (continued)

		G	T	т	С	G	A	Α	C	G	G	G	G	G	С	Т	G	G	G	G	С	Majority	
5						 .					-							- - .			-+-	•	
J			- -			<u> </u>				-	35 -+-				- -	. - -					36 -+-		
•	341																					cloned ac 43030 sl	hc
	338 338	-																				Blast ac shc cloned at 49025 sl	ha
10	338																					Blast at shc	110
		G	C	7)	т	λ	C	æ	7	T	G	т	æ	æ	λ	۳	Δ	Δ	ر	λ	C	Majority	
· · · · · · · · · · · · · · · · · · ·																					-+-	Majority	
1.5											-	70							•		38	0	
15	361		٠	Δ.	·						•										-+-	cloned ac 43030 sl	hc
	358																					Blast ac shc	
	358																					cloned at 49025 sl	hc
20	358	G	C	A	T	Α	С	G	A	T	G	T	G	G	Α	C	A	A	С	A	C	Blast at shc	
20		G	С	G																		Majority	
																					+-	•	
			. – .								35 -+-						. – -				40 -+-	U	
25	381	G	Α	G	C	G	A	Т	C	Т	C	С	C	Ģ	Α	A	_	С	C	A	C	cloned ac 43030 sl	hc
																						Blast ac shc	
	378																					cloned at 49025 sl	hc
	378	G	С	G	Т	С	A	G	T	Т	G.	A	C	С	A	A	A	Т	С	G	G	Blast at shc	
30		A	Т	T	С	С															G +-	Majority	
											41										42)	D	
											41	.0			· -				. .		420		
25	400						G	 Т	т	c	41 +- T	.0	G	 C	G G	 A		T	T	C	420 +- G	cloned ac 43030 sl	hc
35	397	A	T	C	C	C	G G	T T	T T	C	41 +- T T	.0	G G	C C	G G	A A	c c	T T	T T	C	420 +- G G	cloned ac 43030 sl Blast ac shc	
35	397 397	A A	T T	C T	C C	C C	G G A	T T	T T T	CCT	41 T T	.0 - -	G G G	CCC	G G A	A A A	c c	T T	T T		420 +- G G	cloned ac 43030 sl Blast ac shc cloned at 49025 sl	
35	397	A A	T T	C T	C C	C C	G G A	T T	T T T	CCT	41 T T	.0 - -	G G G	CCC	G G A	A A A	c c	T T	T T		420 +- G G	cloned ac 43030 sl Blast ac shc	
	397 397	A A A	T T	C T T	C C C	С С С	G G A	T T T	T T T	CCTT	41 T T	.0 - - - T	G G G	C C C	G G A G	A A A	C C C	T T T	T T T	0000	420 +- G G G G	cloned ac 43030 sl Blast ac shc cloned at 49025 sl	
35 40	397 397	A A A	T T	C T T	C C C	С С С	G G A	T T T	T T T	CCTT	41 T T	.0 - - T	G G G	C C C	G G A G	A A A	C C C	T T T	T T T	0000	420 +- G G G G	cloned ac 43030 sl Blast ac shc cloned at 49025 sl Blast at shc Majority	
	397 397 398	A A A	T T	C T C	C C G	C C A	G G A A	T T T	T T T T	C C T T	41 T T T T A +-	.0 - T T	G G G T	C C C C	G G A G	A A A T	c c c c	T T T	T T T G	0000	420 G G G C +- 440	cloned ac 43030 sl Blast ac shc cloned at 49025 sl Blast at shc Majority	hc
	397 397 398	A A A G	T T	CTTC	C C G	C C A A	G G A A	T T T G	T T T T	C C T T G	41 TTTTT A +- 43	.0 - - T T	G G G T	C C C G	G G A G	A A A T	C C C C	T T T C	T T T G	0000	420 +- G G G C +- 440	cloned ac 43030 sl Blast ac shc cloned at 49025 sl Blast at shc Majority cloned ac 43030 sl	hc
40	397 397 398 419 416	A A G G G G	T T	C T T C C C	C C G G G	C C A A A	G G A A A A A A	T T G G G	T T T T T T T T	CCTTGGG	41 + T T T T A + 4 A A	0 T T C C C	G G G T	C C C C G G	G G A G A	AAAA T	000000000000000000000000000000000000000	T T T C C C	T T T G G G	0000	420 +- G G G G C +- 440 +- C	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc	hc
40	397 397 398 419 416 416	A A G G G G	T T	CTT C	C C C G G G G	C C A A A A	G G A A A A A A	T T T G G G G	T T T T T T T T T	CCTTGGGGG	41- TTTT A + 43 + A A A	0 T T C C T	G G G T C C T	C C C C G G G G	G G A G A A A	A A A T T T	0000	TTT C C C C	T T G G G G	0000	420 G G G C + C C C	cloned ac 43030 sl Blast ac shc cloned at 49025 sl Blast at shc Majority cloned ac 43030 sl	hc
40	397 397 398 419 416 416	A A A G G G G G	T T T	CTTCCCC	C C C G G G G	C C C A A A A A	G G A A A A A A A A	TTTGGGGGG	TTTTTTTTTTT	CCTTGGGGG	4+TTTT A+4+AAAA	O T T T O C C T T	G G G T C C T T	C C C C G G G G	G G A G A A A A A	AAAA T TTTT	0000	TTTCCCCC	T T G G G G G	0000 0	420 +-GGGGGC-+-CCCC	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc	hc
40	397 397 398 419 416 416	A A A G G G G G	T T T	CTT C	C C C G G G G	C C C A A A A A G	G G A A A A A A A	T T T G G G G G A	TTTTTTTTG	CCTT GGGGGA	41- TTTT A + 4 + A A A A C	0 - T T T C C T T G	G G G T C C T T T	C C C C G G G C	G G A G A A A A A	AAAA T TTTT C	בספט ט	TTTT CCCCC	TTTT GGGGG	0000 0 00000	420 +-GGGGGC-+- 440 +-CCCCC	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh	hc
40	397 397 398 419 416 416	A A A G G G G G	T T T	C T T C C C C C	C C C G G G G	C C C A A A A G	G G A A A A A A A A	T T T G G G G A	TTTTT	CCTTGGGGGA	41- TTTT A + 4 + AAAA C + 45	0 - T T O C C T T G O	G G G T C C T T T	C C C C G G G G C	G G A A A A A A	AAAA T TTTT C	0000 0	TTT C CCCC G	TTT G G G G C	0000 0 0000 0	420 +-GGGGGC-+-CCCCC-+-460	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority Majority	hc
40	397 397 398 419 416 416 418	A A A G G G G G G	T T T	CTT C	C C C G G G G	C C C A A A A A G	G G A A A A A A A	T T T G G G G G A	TTTT	CCTTGGGGGA	4+TTTT A+4+AAAA C+4+	0 - T T O C C T T G - 0	G G G T C C T T T	C C C C C G G G G C	G G A G A A A A A	AAAA T TTTT C	0000 0	TTTT C	TTT GGGGG	0000 0 0000 0	420 +-GGGGGC-+-440 -+-CCCCC-+-460	cloned ac 43030 shelast ac shc cloned at 49025 shelast at shc Majority cloned ac 43030 shelast ac shc cloned at 49025 shelast at shc Majority Majority	hc
40	397 398 419 416 416 418	A A A G G G G G G	TTT	CTT C	C C C G G G G G G	C C C A A A A A G G	GGAA A A A A A A	TTTT G GGGG A A	TTTT T	CCTT G GGGG A	41-TTTT A+4-AAAA C+45-	O T T O CCTT G O G	GGGG T CCTT T		G G A G A A A A A	AAAA T TTTT C C	בסטטט טייים מטטטט טיייים	TTTT C CCCC G	TTTT G GGGG C	0000 0 0000 0 0	420 G G G C +-440 C C C C C +-60 C +-C	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned at 49025 sh Cloned at 49025 sh Cloned at 49025 sh Cloned ac 43030 sh	hc
40	397 398 419 416 416 418	A A A G G G G G G G G G	TTT	C T T C C C C C C C C C C C C C C C C C	C C C G G G G G A	C C C A A A A A G G G	G G A A A A A A A A	TTTT G G G G G A G	TTTT T G G G	CCTT G GGGG A A A	41-TTTT A + 43 + AAAA C + 4 + C C	O T T O CCTT G O G	GGGG T CCTT T		G G A A A A A A A	AAAA T TTTT C C C C	מטטט טייים מטטט טייים ט	TTTT C CCCC G G	TTTT G GGGG C C CC		420 +-GGGGGC++40 +-CCCCCCC+-60 +-CCCCCCCCC	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc	hc hc
40	397 398 419 416 416 418	A A A G G G G G G A	TTT		C C C G G G G G A G	C C C A A A A A G G G G	G G A A A A A A A A A A A A A A A A A A	TTTT G G G G G A A G A	TTTT T G G G G	CCTT G GGGG A A A A	41 TTTT A + 4 + A A A A C + 4 + C C C	O T T O CCTT G O G G G	GGGG T CCTT T TTT	C C C C G - G G G G C - C C C	G G A G A A A A A A A A A A A A A A A A	AAAA T TTTT C CCC		TTTT C CCCC G GGG	TTTT G GGGG C C CCC	CCCC C CCCA	420 +-GGGGGC+440 +-CCCCCCC+60	cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned ac 43030 sh Blast ac shc cloned at 49025 sh Blast at shc Majority cloned at 49025 sh Cloned at 49025 sh Cloned at 49025 sh Cloned ac 43030 sh	hc hc

Figure 5: She polynucleotide sequence alignments (continued)

			0											_			_	
			G T														Majority	
5							47	0								480		
	455	A C A C	G T	G C	T	C 9	A	G G	T (G 7	r 1 r 1	' T	C	G G	G G	<u>C</u>	cloned ac 43030 Blast ac shc	
10	455 458																cloned at 49025 Blast at shc	shc
		A G	СТ	T 7		G									G 		Majority	
1.5					•		49	Ó		•			·			500)·	
15	475	A G A G	C T	TO	C G	G (T	A A	C	G 2 G 2	A C	: G	A	C C	G G	C C	cloned ac 43030 Blast ac shc cloned at 49025	
20																	Blast at shc	
20		СТ	G G	A Z	A G	G :	G	A	T	T (C 0	; G	C	G	G	G ı	Majority	
							5	0								520)	
25	495	C T	G G	AA	A G	G :	G	A A	T T	C (C A	A G	C	G G	C C	G G	cloned ac 43030 Blast ac shc cloned at 49025	
	498	AT	G G	A	A G	G :	G	A	T.	T (0 0	3 0	: A	Α	G	G	Blast at shc	
30			G I														Majority	
							5.	3 0								54		
35	515	C G C G	G T G T	. G (3 C 3 A 3 A	G 'A '	r A r A r A	C T	С С	T (T (C <i>I</i> C <i>I</i>	A A A A	G G G	C C G	G G C	G G G	cloned ac 43030 Blast ac shc cloned at 49025	
	518																Blast at shc	
40		G A	G C	A (3 C		3 C										Majority	
								50 					. 			56	0	
45	535 535	G A	G C	ACAC	G A G A A C	A G	3 C	C C C	G G A	G. G.	A (A (A :		6 G 6 G 6 G	C G	A A T	G G C	cloned ac 43030 Blast ac shc cloned at 49025 Blast at shc	
		СТ	G G														Majority	
50				· -			5	70								58	0	
	558 555 555	CI	G	T	T C	G	G T	C C	G G	C T	T (3 (3	3 G	G	C	.G G	cloned ac 43030 Blast ac shc cloned at 49025	
55	558																Blast at shc	

Figure 5: Shc polynucleotide sequence alignments (continued)

		CAACTACGTGTATGGC	
5		590	600
10	578 575 575 578	C A A C T A C A T C T A C G G C C A A T T A C C T C T A C G G C C A A C T A C G T G T A T G G C C A A C T A C G T G T A T G G C	A C G cloned ac 43030 shc A C G Blast ac shc A T C cloned at 49025 shc
		G C G C G G T G G T T T C G G G	
1.5		610	620
15 20	598 595 595 598	G C G C G G T G G T G T C G G C G C G G T G T	G C T cloned ac 43030 shc G C T Blast ac shc A C T cloned at 49025 shc
20		A A G G C G G T C G G T C T C G .	•
		630	640
25		A A G G C G G T C G G G A T C G . A A G G C G G T C G G G A T C G . A A G G C C G T C G G T G T C G . A A G G C C G T C G G T G T C G .	A C A cloned ac 43030 shc A C A Blast ac shc A T A cloned at 49025 shc
30		GCGTGAGCCGTGGGTT	3 . 1
		650	660
35	638 635 635 638	G C G C G A G C C G T A C A T T C G C G C G A G C C G T A C A T T C G C G T G A G C C G T G G G T G G C G T G A G C C G T G G G T G G C G T G A G C C G T G G G T G	C A A cloned ac 43030 shc C A A Blast ac shc C A A cloned at 49025 shc
40		A G T C G C T C G A C T G G G T (
.0		670	680
45	655	A G G C G C T C G A T T G G G T C A G G C G C T C G A C T G G G T C A G T C G C T C G A C T G G C T C A G T C G C T C G A C T G G C T	G G A cloned ac 43030 shc C G A Blast ac shc C G T cloned at 49025 shc
		G A G C A T C A G A A T G C G G	
50		690	700
55	675 675	C A G C A T C A G A A C C C G G A C A G C A T C A G A A C C C G G A G A G C A T C A A A A T G A G G A G A G C A T C A A A A T G A G G A	A C G cloned ac 43030 shc A C G Blast ac shc A T G cloned at 49025 shc

Figure 5: Shc polynucleotide sequence alignments (continued)

	•	GCGGCTGGGTGAAGACTG - Majority
5		710 720
10	698 695 695 698	G CGGCTGGGGTGAAAGCCGA Cloned at 43030 shc
		CCGXTCXTACGAGGATCC Majority
1.0		730 740
15 20	698 714 715 717	
		G X X X C T C G C G G G T C A G G G C G Majority
		750 760
25	698 732 735 735	
30		CGAGXACACCGTCGCAGACX Majority
		770 780
35	698 752 755 755	Cloned ac 43030 shc CGAGCACCCGTCGCAGACG Blast ac shc CGGTACCAGACCGTAA CCAACCGTCGCAGACC Blast at shc
40		GCCTGGGCGTTGATGGCGCT Majority
70		790 800
45	698 772 775 775	Cloned ac 43030 shc G C C T G G G C C T G A T G G C G C T T C A T G G T C A T A G C T G T T T C C G C C T G G G C G T T G A T G G C G C T Blast ac shc G C C T G G G C G T T G A T G G C G C T Blast at shc
		CATCGCGGGCGGCXGTGTCG Majority
50		810 820
55	698 792 795 795	Cloned ac 43030 shc CATCGCGGGCCGCCAGGCAGGGGGGGGGGGGGGGGGGG

Figure 5: Shc polynucleotide sequence alignments (continued)

																						Majority
5											83										+- 84	D
				•							.+-										+-	
	698																					cloned ac 43030 shc
	812_	Α	G	Т	С	С	G	Α	G	G	C	C	G	C	G	C	G	C	С	G	C	Blast ac shc
																						cloned at 49025 shc
10	815	A	G	T	C	A	G	A	T	G	.C	G	G	Т	Α	T	Т	G	С	G	C	Blast at shc
		G	G	G	G	т	C	С	х	х	т	Α	C	C	т	х	X	х	х	G	_	Majority
,																					-+-	-
	•										85										86)
15											+-										-+-	
	698																					cloned ac 43030 shc
																						Blast ac shc
																						cloned at 49025 shc
	835	G	Ġ	G	G	T	C	Α	С	T	Т	A	С	C	T	\mathbf{T}	С	A	C	G	-	Blast at shc
20																						-10- • • • ·
																						Majority
											•										-+-	_
		 – .	87				 .						88 -+-	
25	698										•										•	cloned ac 43030 shc
	851	Ά	G	Α	C	G	C	A	G	С	G	С	С	C	G	G	Α	С	G	_	G	Blast ac shc
																						cloned at 49025 shc
																						Blast at shc
•			•		_		_															
30		т	G	G	С	T	G	х	Х	х												Majority
											-											
	600										-											cloned ac 43030 shc
35	698 870	_	~	~	~	m	c	C	a	7												Blast ac shc
<i>)</i>	870 873							G	G	A												cloned at 49025 shc
		T																				Blast at shc
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Figure 6: She amino acid sequence alignments

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10	1																		L				A.acidoterrestris DSM 3902
	1																		-				Bacillus subtilis
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	9	_																					A. acidocaldarius JCM 5260T
	9	-																					A.acidoterrestris DSM 3902 Bacillus subtilis
25	4 21																						Dictyostelium discoideum
23	7					_													-				Synechocystis sp. PCC 6803
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35	10																					A	A.acidoterrestris DSM 3902
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	41																		T				Dictyostelium discoideum
	7																		Q				Synechocystis sp. PCC 6803
40	28	-		Ρ	A	A	A	A	G	٧	-	-	-	-	•	-	Ъ	ĸ	A	Α	А	R	Streptomyces coelicolor A3
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	14																		G				A.acidoterrestris DSM 3902
	11																		G				Bacillus subtilis
	61																					A	Dictyostelium discoideum
50	20																		G				Synechocystis sp. PCC 6803 Streptomyces coelicolor A3
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Figure 6: Shc amino acid sequence alignments (continued)

GPLLSNVTMEAEYVLLCHIL Majority ------5 90 ______ 34 GPLLSNVTMEAEYVLLCHIL A. acidocaldarius ATCC27009 34 GPLLSNVTMEAEYVLLCHIL A. acidocaldarius JCM 5260T 34 APLLSNVCMEAEYVLLCHCL A.acidoterrestris DSM 3902 10 31 FCFEGPIMINSFFILLLTSL Bacillus subtilis 81 GDYGGPMFLLPGLVITCYVT Dictyostelium discoideum 40 SELESNVTITAEVVILHKIW Synechocystis sp. PCC 6803 61 GDLETNVTMDAEDLLLRQFL Streptomyces coelicolor A3 GRVDRER - - MEKIRRYLLHE Majority 15 _____+ 110 ______ 54 DRVDRDR - - MEKIRRYLLHE A. acidocaldarius ATCC27009 54 DRVDRDR - - MEKIRRYLLHE A. acidocaldarius JCM 5260T 20 54 GKKNPER - - EAQIRKYIISQ A.acidoterrestris DSM 3902 51 DEGENEKELISSLAAGIHAK Bacillus subtilis 101 GYQLPESTQREIIRYLFNRQ Dictyostelium discoideum 60 GTAAQRP - LEKAKNYLLQQ Synechocystis sp. PCC 6803 81 GIQDEET - - TRAAALFIRGE Streptomyces coelicolor A3 25 QREDGTWALYPGGP-GDLST Majority ______ 30 QREDGTWALYPGGP-PDLDT A. acidocaldarius ATCC27009 72 QREDGTWALYPGGP-PDLDT A. acidocaldarius JCM 5260T 72 RREDGTWSIYPGGP-SDLNA A.acidoterrestris DSM 3902 71 QQPDGTFINYPDETRGNLTA Bacillus subtilis 35 121 NPVDGGWGLHIEAHSDIFGT Dictyostelium discoideum 78 QRDHGGWELYYGDG-GELST Synechocystis sp. PCC 6803 99 QREDGTWATFYGGP-GELST Streptomyces coelicolor A3 TVEAYVALKYLG-VSADEPH Majority _____+-40 150 -------TIEAYVALKYIG - MSRDEEP A. acidocaldarius ATCC27009 91 TIEAYVALKYIG - MSRDEEP A. acidocaldarius JCM 5260T 91 TVEAYVALKYLG - EPASDPQ A.acidoterrestris DSM 3902 45 91 TVQGYVGMLASGCFHRTEPH Bacillus subtilis 141 TLQ-YVSLRLLG-VPADHPS Dictyostelium discoideum 97 SVEAYTALRILG - VPATDPA Synechocystis sp. PCC 6803 118 TIEAYVALRLAG - DSPEAPH Streptomyces coelicolor A3

Figure 6: Shc amino acid sequence alignments (continued)

		M V K A L E F I Q S Q G G I E S S R V F Majority
5		170 180
10	110 110 110 111 159 116 137	M Q K A L R F I Q S Q G G I E S S R V F A. acidocaldarius ATCC27009 M Q K A L R F I Q S Q G G I E S S R V F A. acidocaldarius JCM 5260T M V Q A K E F I Q N E G G I E S T R V F A.acidoterrestris DSM 3902 M K K A E Q F I I S H G G L R H V H F M Bacillus subtilis V V K A R T F L L Q N G G A T G I P S W Dictyostelium discoideum L V K A K N F I V G R G G I S K S R I F Synechocystis sp. PCC 6803 M A R A A E W I R S R G G I A S A R V F Streptomyces coelicolor A3
15		TRMWLALVGEYPWDKLPMIP Majority
		190 200
20	130 130 130 131 179	TRMWLALVGEYPWEKVPMVP A. acidocaldarius ATCC27009 TRMWLALVGEYPWEKVPMVP A. acidocaldarius JCM 5260T TRLWLAMVGQYPWDKLPVIP A.acidoterrestris DSM 3902 TKWMLAANGLYPWPAL-YLP Bacillus subtilis GKFWLATLNAYDWNGLNPIP Dictyostelium discoideum
25	136 157	T K M H L A L I G C Y D W R G T P S I P Synechocystis sp. PCC 6803 T R I W L A L F G W W K W D D L P E L P Streptomyces coelicolor A3
		PEIMLLPKNVPLNIYEFGSW Majority
30		210 220
35 .	150 150 150 150 199 156 177	PEIMFLGKRMPLNIYEFGSW A. acidocaldarius ATCC27009 PEIMFLGKRMPLNIYEFGSW A. acidocaldarius JCM 5260T PEIMHLPKSVPLNIYDFASW A.acidoterrestris DSM 3902 LSLMALPPTLPIHFYQFSSY Bacillus subtilis IEFWLLPYNLPIAPGRWWCH Dictyostelium discoideum PWVMLLPNNFFFNIYEMSSW Synechocystis sp. PCC 6803 PELIYFPTWVPLNIYDFGCW Streptomyces coelicolor A3
40		ARATVVPLSIVMAQQPV Majority
		230 240
45	170 170 170 170 219 176 197	A R A T V V A L S I V M S R Q P V A. acidocaldarius ATCC27009 A R A T V V A L S I V M S R Q P V A. acidocaldarius JCM 5260T A R A T I V T L S Y R H E S P T C A.acidoterrestris DSM 3902 A R I H F A P M A V T L N Q R Bacillus subtilis C R M V Y L P M S Y I Y A K K T T G P L Dictyostelium discoideum A R S S T V P L M I V C D Q K P V Synechocystis sp. PCC 6803 A R Q T I V P L T I V S A K R P V R P A Streptomyces coelicolor A3

Figure 6: Shc amino acid sequence alignments (continued)

		_	F	P	L	P	E	L	Α	R	V	₽	E	L	Y	E	Т	D	V	P	P	Majority
5											+- 25		•								+- 26	0
3																. - -					+-	
	187																					A. acidocaldarius ATCC27009
	187	_																				A. acidocaldarius JCM 5260T
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10	185	_	r R	v	т.	т	Ŋ	R	_	N	I	s	s	L	H	Н	L	D	_	_		Bacillus subtilis
10	239	т	D.	т.	v	ĸ	D	T.	R	R	_	_	E	I	Y	C	0	E	Y	E	K	Dictyostelium discoideum
	193	_																				Synechocystis sp. PCC 6803
	217	P	F	Р.	L	D	Ē	L	H	T	D	P	Α	-	-	_	R	P	N	P	P	Streptomyces coelicolor A3
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	206	R	R	_	R	G	Α	к	G	G	G	G	W	_	_	_	I	F	D	A	_	A. acidocaldarius ATCC27009
20	206	R	R	_	R	G	Α	K	G	G	G	G	W	-	_	-	I	F	D	Α	-	A. acidocaldarius JCM 5260T
-	206	K	R	_	R	S	A	K	G	G	D	S	G	-	_		F	F	ν	Α	-	A.acidoterrestris DSM 3902
	201	H	M	Т	K	N	P	F	T	W	L	R	s	_	-	D	A	F	Ε	E	R	Bacillus subtilis
	257	I	N	W	s	E	Q	R	N	N	Ι	s	K	L	D	М	Y	Y	E	H	T	Dictyostelium discoideum
	212	V	Q	Y	K	L	₽	E	S	G	T	I	W	-	-	D	I	F	I	G	-	Synechocystis sp. PCC 6803
25	234	R	P	-	L	A	P	V	A	S	W	D	G	-	-	-	A	F	Q	R	-	Streptomyces coelicolor A3
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	221		L	D	R	A	ь	H	G	Y	Q	K	L	-	-	s	٧	Н	P	F	R	A. acidocaldarius ATCC27009
	221	-	L	D	R	Α	L	H	G	Y	Q	K	L	-	-	S	V	H	P	F	R	A. acidocaldarius JCM 5260T
	221																				R	A.acidoterrestris DSM 3902
	219																				Q	Bacillus subtilis
35	277	S	L	L	N	V	I	N	G	S	L	N	A	Y	E	K	V	H	.S	K	W	Dictyostelium discoideum
	229	-	L	D	S	L	F	K	L	Q	E	Q	A	-	-	K	V	V	₽	F	R	Synechocystis sp. PCC 6803
	249	-	I	D	K	A	L	H	A	Y	R	K	V	-	-	A.	P	R	R	L	R	Streptomyces coelicolor A3
		R	A:	G	E	Α	R	A	ь	т	W	I	L	E	R	Q	E	G	D	G	S	Majority
40											-+-					- -					-+-	•
	·										_										32	
																					-+-	A. acidocaldarius ATCC27009
	238																				S	A. acidocaldarius JCM 5260T
15	238						R														C	A.acidoterrestris DSM 3902
45	238	K	5	ن	E T	Š	T.	A	A. Tì	T.	v	M	T.	ת	D	Z.	中	K	ם	G	T	Bacillus subtilis
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	246	r.		5	T.	7	T.	Δ	E	ĸ	W	Т	Ţ.	E	R	ō	E	v	s	G	D	Synechocystis sp. PCC 6803
	266																				C	Streptomyces coelicolor A3

Figure 6: Shc amino acid sequence alignments (continued)

	,	W G G I Q P P W F Y A L I A L K V L G M Majority	
5		330 340	
10	258 258 258 259 317 266 286	W G G I Q P P W F Y A L I A L K I L D M A. acidocald W G G I Q P P W F Y A L I A L K I L D M A. acidocald W G G I Q P P W F Y A L L A L K C L N M A.acidoterrolly S Y A S A T I Y M V Y S L L S L G V Bacillus sulk Y I D I G P V N K T V N M L C V W D R Dictyostelium G G I I P A M L N S L L A L K V L G Y Synechocyst	darius ATCC27009 darius JCM 5260T estris DSM 3902 btilis um discoideum is sp. PCC 6803 s coelicolor A3
15		T - Q H P A F I K G L E G L E L Y G V E Majority	
		350 360	•
20	278 278 278 279 337	T - Q H P A F I K G W E G L E L Y G V E A. acidocalor - Q H P A F I K G W E G L E L Y G V E A. acidocalor - D H P A F V K G F E G L E A Y G V H A.acidoterror R Y S P I I R R A I T G I K S L V T K Bacillus su	darius JCM 5260T estris DSM 3902
25	286 306		is sp. PCC 6803 s coelicolor A3
		LSDGGWMFQA-SISPVWDTG Majority	
30		370 380	
30	297 297	L D Y G G W M F Q A - S I S P V W D T G A. acidocal L D Y G G W M F Q A - S I S P V W D T G A. acidocal T S D G G W M F Q A - S I S P I W D T G A.acidoterr	darius JCM 5260T estris DSM 3902
35	299 356 306 326	C N G I P Y L - E N - S T S T V W D T A Bacillus su L S F D G M K M Q G Y N G S Q L W D T A Dictyosteli T E - D S Y A I Q A - C V S P V W D T A Synechocyst R E D G A R M I E A - C Q S P V W D T C Streptomyce	um discoideum
40		LAVLALRAAGLPADHPALVK Majority	
	216	+-	darius ATCC27009
45	316 316 317 376	LAVLALRAAGLPADHDRLVK A. acidocal LTVLALRSAGLPPDHPALIK A.acidoterr LISYALQKNGVTETDGSVTK Bacillus su FTIQAFMESGIANQFQDCMK Dictyosteli	darius JCM 5260T estris DSM 3902
50	324 345		s coelicolor A3
50			

Figure 6: Shc amino acid sequence alignments (continued)

		GEWLLDRQITVPGDWAVKR	
5			20
10	336 336 336 337 396 344 365	G E W L L D R Q I T V P G D W A V K R G E W L L D R Q I T V P G D W A V K R G E W L V S K Q I L K D G D W K V R R A D F L L E R Q H T K I A D W S V K N A G H Y L D I S Q V P E D A R D M K H G Q W L L D K Q I L T Y G D W Q I K N A S D W M L G E Q I V R P G D W S V K R	A. acidocaldarius ATCC27009 A. acidocaldarius JCM 5260T A.acidoterrestris DSM 3902 Bacillus subtilis Dictyostelium discoideum Synechocystis sp. PCC 6803
15		- PNLKPGGWAFEFDNVNYP	
			40
20	356 356 356 357 416 364 385	- P N L K P G G F A F Q F D N V Y Y F - P N L K P G G F A F Q F D N V Y Y F - R K A K P G G W A F E F H C E N Y F - P N S V P G G W G F S N I N T N N F H R H Y S K G A W P F S T V D H G W F - P H G E P G A W A F E F D N N F Y F - P G L P P G G W A F E F H N D N Y F	A. acidocaldarius ATCC27009 A. acidocaldarius JCM 5260T A.acidoterrestris DSM 3902 Bacillus subtilis Dictyostelium discoideum Synechocystis sp. PCC 6803
		OVDDTAVVVLALNGLRI	Majority
30		450 4 	60
35	374 374 375 436 382 403	O V D D T A V V V W A L N T L R I O V D D T A V V V W A L N T L R I O V D D T A M V V L A L N G I Q I O C D D T T A V L K A I P R N H S I S D C T A E G I K S A L A L R S L P E O I D D T C V V M M A L Q G I T I O I D D T A E V V L A L R R V R E	A. acidocaldarius JCM 5260T A.acidoterrestris DSM 3902 Bacillus subtilis Dictyostelium discoideum Synechocystis sp. PCC 6803
40		PDEERRRDAITKGFRWLLGM	· -
45	391 391 391 392 456 399 420	470 P D E R R R R D A M T K G F R W I V G M P D E R R R R D A M T K G F R W I V G M P D E G K R R D A L T R G F R W L R E M P A A W E R G V S W L L S M I E P I S L D R - I A D G I N V L L T I P D E E R K Q G A I N K A L Q W I A T M H D P E R V E K A I G R G V R W N L G M	A. acidocaldarius ATCC27009 A. acidocaldarius JCM 5260T A.acidoterrestris DSM 3902 Bacillus subtilis Dictyostelium discoideum Synechocystis sp. PCC 6803

Figure 6: Shc amino acid sequence alignments (continued)

		QSSNGGWGAYDVDNTSDLPN Majority	٠.
5		490 500	
10	411 411 411 406 475	Q S S N G G W G A Y D V D N T S D L P N A. acidocal Q S S N G G W G A Y D V D N T S D L P N A. acidocal Q S S N G G W G A Y D V D N T R Q L T K A.acidoterr Q N N D G G F S A F E K N V N H P L I R Bacillus su	darius JCM 5260T estris DSM 3902 btilis
15	419 440	QCKTGGWAAFDIDNDQDWLN Synechocyst	um discoideum is sp. PCC 6803 s coelicolor A3
		510 520	•
20	431 431 431 426 495 439	H I P - F C D F G E V - T D P P S E D V A. acidocal S D S I F A T S G E V - I D P P S E D V A.acidoterr L L P L E S A E D A A - V D P S T A D L Bacillus su K F N P S E V F Q N I M I D Y S Y V E C Dictyosteli	darius ATCC27009 darius JCM 5260T estris DSM 3902 btilis um discoideum is sp. PCC 6803
25	460	•	s coelicolor A3
	•	530 540	
30	449 449 450 445 515	TAHVLECFGSFGA. acidocal TAHVLECFGSFGA. acidocal TAHVLECFGSFGA. acidocal TAHVLECFGSFGBacillus sussa ACIQAMSAFRKHAPNHPR Dictyosteli	darius JCM 5260T estris DSM 3902 btilis
35	457 478	TARVVEMLGACG Synechocyst	is sp. PCC 6803 s coelicolor A3
		Y D E A W K V I R R A V E Y L K R E Q E Majority	
40		550 560	
45	461 462 458 535 469 490	Y D D A W K V I R R A V E Y L K R E Q K A acidocal Y D E A W K A V E Y L K R E Q K A acidocal Y D E A W K R E Q K A acidocal Y D E A W K F I K A acidocal Y D E A W K Y I A acidocal Y D E A W K Y I K A A acidocal Y D E A W K Y I K A A A A A I I I A A A I I I I I	darius JCM 5260T estris DSM 3902

Figure 6: Shc amino acid sequence alignments (continued)

		Q D G S W F G R W G V N Y L Y		
5		570	580	
10	481 481 482 478 555 489 509	P D G S W F G R W G V N Y L Y P D G S W F G R W G V N Y L Y P D G S W F G R W G V N Y V Y Q N G S W Y G R W G V C Y I Y Q D G S W L G S W G I C F T Y Q D G S W F G R W G V N Y L Y T D G S W F G R W G V N Y V Y	Y G T G A V A. acidocaldarius ATCC27009 Y G T G A V A. acidocaldarius JCM 52607 Y G I G A V A.acidoterrestris DSM 3902 Y G T W A A Bacillus subtilis Y G T W F G Dictyostelium discoideum Y G T S G A Synechocystis sp. PCC 6803	
15		V S A L K A V G L D T R E P Y		
		590	600	
20	501 502 498 575	V S A L K A V G I D T R E P V V P G L K A V G V D M R E P V	Y I Q K A L A. acidocaldarius ATCC27009 Y I Q K A L A. acidocaldarius JCM 52607 W V Q K S L A.acidoterrestris DSM 3902 V Y K R L C Bacillus subtilis S I V K A C Dictyostelium discoideum Q I K T A I Synechocystis sp. PCC 6803	
23	329	DWLESHQNADGGWGE	E D C R S Y Majority	
		610	620	
35	521 521 522 518 595 529 549	D W V E Q H Q N P D G G W G E D W L V E H Q N E D G G W G E V	E D C R S Y A. acidocaldarius ATCC27009 E D C R S Y A. acidocaldarius JCM 52607 E D C R S Y A.acidoterrestris DSM 3902 - G S N P Y Bacillus subtilis E S F K S - Dictyostelium discoideum E T C E S Y Synechocystis sp. PCC 6803	
40		E - D P E Y A G Q G A S T A S	· ·	
		630	640	
45	541 541 542 524 614 549 569	E - D P A Y A G K G A S T P S E - D P A Y A G K G A S T P S D - D P R L A G Q G V S T P S K M M T E - A G E N P A K A F N V T K E Y V Q H E T S Q V V K - N K Q L K G Q G N S T A S R Y V R E W S G R G A S T A S	S Q T A W A A. acidocaldarius ATCC27009 S Q T A W A A. acidocaldarius JCM 52607 S Q T A W A A.acidoterrestris DSM 3902 P K S K Bacillus subtilis V N T G W A Dictyostelium discoideum S Q T A W A Synechocystis sp. PCC 6803	

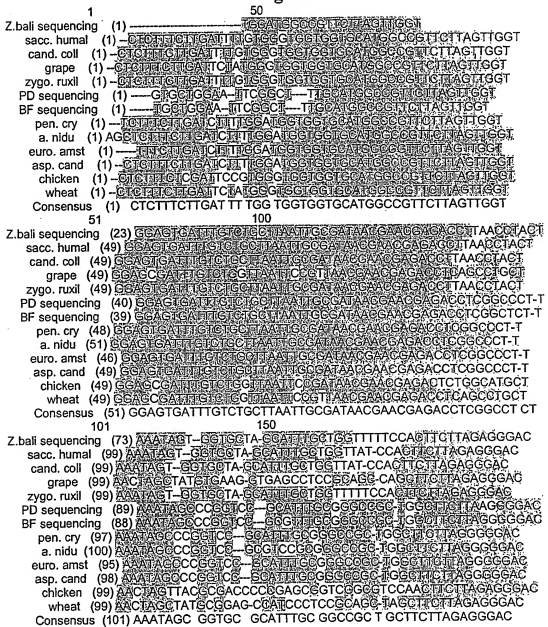
Figure 6: She amino acid sequence alignments (continued)

		LMALIAG GRAE Majority	
5		650 660	
10	560 561 541 634 568 589	L M A L I A G G R A E A. acidocaldarius A' L M A L I A G G R A E A. acidocaldarius JG L M A L I A G G R V E A.acidoterrestris DG Bacillus subtilis L L S L M S A - K Y P D R Dictyostelium discos L I G L L D A L K Y L P S L G Q D A K L Synechocystis sp. PG L M A L L A A G E R D - Streptomyces coelication	CM 5260T SM 3902 ideum CC 6803
15		SEAAERGVAYLVETQRPDGG Majority	٠
		670 680	
20	571 571 572 541	S E A A R R G V Q Y L V E T Q R P D G G A. acidocaldarius A S E A A R R G V Q Y L V E T Q R P D G G A. acidocaldarius J G S D A V L R G V T Y L H D T Q R A D G G A. acidoterrestris D G C C C C C C C C C C C C C C C C C C	CM 5260T
25	646 588 600	- E C I E R G I K F L I Q R Q Y P N G D Dictyostelium discort T A I E G G V A F L V Q G Q T P K G T Synechocystis sp. PG S K A V E R G V A W L A A T Q R E D G S Streptomyces coelicord	CC 6803
		WDEPYYTGTGFPGDFYLGYT Majority	
30	•	690 700	
35	591 591 592 541 665 608 620	W D E P Y Y T G T A S P G D F Y L G Y T A. acidocaldarius AT W D E P Y Y T G T G F P G D F Y L G Y T A. acidocaldarius J G W D E E V Y T G T G F P G D F Y L A Y T A.acidoterrestris DS	CM 5260T SM 3902 ideum CC 6803
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Figure 6: Shc amino acid sequence alignments (continued)

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GTGGTGCTAGCATTTGCTG Yeast prime up CCGCTGGCTTCTTAGGG Mold prime up

Figure 7 (continued)

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Z.bali sequencing (121) TATCGGTU	CAAGCCGATGGAACHTGACGCAATAACAGGTGTGTGATGC
sacc. humal (145) TATCGGTTT	AAGDCGATGGAAGTTTCAGGGATAACAGGTCTGTCATGC AGCCGATGGAAGTTTGAGGCAATAACAGGTCTGTCATGC
cand. coll (145) TATOGGT TEA	ACCGATGGAAGTITGACGGAATAACAGGTCTGTGATGG
grape (147) TATIGGCOGCTT	accicaaccaac iiiicacccaaiaacaccicicicatcc
zygo. ruxil (146) TATEGETITEA	AGCCGATGGAAGITHGAGGCAATAACAGGTOTGTGATGC
PD sequencing (135) TATEGES	-CAACCCGATCGAACTGCCCCSGSAATAACACCHD ICHGATGC
BF sequencing (134) TATGGGGT	CAAGCCGATGGAAGTGCGCGGCAATAACAGGTCTGTAATGC
pen. cry (143) TATEGGGT-CA	AGCGGATIGGAAGT GCGCGGCAATAACAGGTIGTGTGATGC
a. nidu (146) TATCGGGT-CA	AGGGGATGGAAGTGCECGGGAANAACAGGTUTGTGATGC
euro. amst (141) TATCGGCT-C	AGGCGATGGAAGTGCGCGGGAATAACAGGTGTGTGTGATGC
asp. cand (144) TATEGECT-O	VAGCCGATGGAAGTIGCGCGGCAATLAACAGGTCTGTGATGC CAGGCACCCGAGATTGAG-CAATLAACAGGTCTGTGATGC
chicken (149) AAGTGGCGTT	CAGEG ACCCGAGATEGAG-GAATAACAGGICIGIGAIGC
wheat (147) TATESCOGTTT	AGGEGACEGAAETHICAGE CAATAACAGGTGTGTGATGC
	CAAGCCGATGGAAGTTTGAGGCAATAACAGGTCTGTGATGC
201	250
Z.bali sequencing (171) CG1 AGAC	GIIII GIIGGGGGGCGCGGGGGGAGCCAGCGAGT
sacc. humal (195) GENAGACE	HIGTGGGCGCACGGGGGGTACAGTGACGGAGCCAGCGAGT
cand. coll (195) GC WAGACG	GIEGEGEGACEGEGETACATTGACGGAGCCAGCGAGT
grape (197) COMAGATON	OTEGGCCGCAEGCGCGACACTGATGTATTCAACGAGT
zygo. ruxil (196) @@ijinGACGii	TETEGEEEGEAGGEGEGTACAETGAGGGAGCEAACGAGT
PD sequencing (184) @GHUAGAU	ethetegecegeacecectavactgacasescaccact
BF sequencing (183) GCINAGAD	STITET GGGGGGGAGGGGGGGGGGGGGGGGGGGGGGGGGGGG
pen. cry (192) CGTTAGATGTL	OTGGGCGCACGOGOGOTACACTGACAGGGGCCAGCGAGT
a. nidu (195) CENIAGAIGH	TIGEGGGGCACGGGCGAGTACACTGACAGGGCCAGCGAGT
euro. amst (190) COMAGANG	TOTEGGECGCACECECETACACTGACAGECCAGCGAGT
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chicken (197) GERLAGARGIN	FPTTOPACTOPETORALINATION OF THE PROPERTY OF TH
wheat (197) @ 124541611	CTGGGCCGACGOGCGTACACTGATGTATTCAACGAGT
	TTCTGGGCCGCACGCGCGCTACACTGAC GGGCCAGCGAGT
251	300 IGGCCBACAGCICIGGCTAATCITGIGAAACTCCGTCCIGC
Z.baii sequencing (221) City—Accura	ECCEACAGETOTTEGTATETTETGAAACTOCGTCGTCC
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cand. coll (245) CHA-Accorded	SCEGAGAGGTCTGGGTAATICTTGTGAAACTCCGTCGTGC GCGGACAGGCCCGGGTAATICTTTG-AAATTTCATCGTGA
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zygo, ruxii (246) CiiA-ACCIII Gi	CCGAGAGAGETCTGGGTAATCTTGTGAAACTCCGTCGTGC TTAACCGAGAGGTTTGGGTAATCTTGTTAAACCCTGTCGTGC
PD sequencing (234) ACAGCAGC	ITIGECCEAEAGETICIGEGTAATOFICTTAAACCCTGTCGTGC
BF sequencing (233) ACAMCAGE	IN ICONOCIONAL CONTROLLA PROPERTINA DE LA CONTROLLA PARA CONTROLLA
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euro, amst (240) ACA CACCA	AACCGAGAGGTCTTGGGTAATCTTGTTAAACCCTGTCGTGC
asp. cand (243) ACATCAGE	GGCGGAGAGGTOTGGGAAAACCCTGTCGTCA
chicken (247) GICIACOGIA	CEGCEGCAGECGCETAACCCCTTIGAACCCCATTCGTGA
wheat (247) ANAMAGEOMIC	GCCACAGCCCCCCTAATCTTGGAAATTTCATCGTGA
Consensus (251) ATAT ACCTT	GGCCGAGAGGTCTGGGTAATCTTGT AAACCC GTCGTGC

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Figure 7 (continued)

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Z.bali sequencing (269) TGGGG	TAGAGCATTGTAATTATTGCTCTTCAACGAGGAATTCCTAGT	AΑ
sacc. humal (293) TSGGGA	AGAGCATTIGTAATTATTGGTCTTCAACGAGGAATTCCTAGTA	A
cand. coll (293) TGGGGAT	BAGGATHGTAATHAHIGGTOINGAAUGAGGAAN TOOTAGHAA	
grape (296) TGGGGATA		
zygo. ruxil (294) IGGGGAT	SAGEATTGTAATTATTGGTCTTCAAGGAGGAATTCCTAGTAA	er 1875.
PD sequencing (284) ISSS	FAGAGCATIGEAATHATIGOTOTICAAOGAGGAATGCCTAG	TAG
BF sequencing (283) TGGGC	HAGAGCATHGCAATHATTGCTCTTCAACGAGGAATGCCTAG HAGAGCATTGCAATTATTGCTCTTCAACGAGGAATGCCTAG	TAG
pen. crv (292) TGGGGAT	SAGCATA GEAATTATA GULU INCAAL GAGGAA LGOGTAGTAG	j
a. nidu (295) IGGGGATA	AGCATTGCAATTATTIGCTETTCAACGAGGAATGCGTAGTAG	
euro. amst (290) TGGGGA	NGAGCATHGGAATHATHGGIGHTCAACGAGGAATIGCCTAGTA	G
asp. cand (293) IGGGGA	GAGCATTGCAATTATTGCTCTTGAAGGAGGAATGCCTAGTA	٥
chicken (297) TGGGGAT	GGGGATIGGAATIATICCCCATGAACGAGGAATITGCCAGTA	A
wheat (297) IGGGGA	ATCATHCAAHGHEGICTICAACGAGGAAHGGCLACTAA	Α.
	AGAGCATTGCAATTATTGCTCTTCAACGAGGAATGCCTAGTA	A
351	GTCATCAACTITECETTGATTACGTCECTGCCCTTTGTACAC	NGΛ
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wheat (347) GCGCGAG	CATCAGOTOGOGTT CACTAGGT COCTGG COTHTTGTACACAC	C
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sacc. humal (393) [SCCSSS]	GCTAG	
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Align	Alignment Report of Ali 16S alignement.meg ClustalV (Weighted)  Thursday, September 04, 2003 10:51 AM	Page 13
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	GAConsensus # GAGGACAGAG Majority	
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Page 1/		307 3 U349	5307 5 U349
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1006 GGGGTCCTTCGGGGCAGAGGACAGG 43030 16s
GGGGTCCCTTCGGGGCAGAGGAGACAGG
GGTTTCCCTTCGGGGCAGGGGAGACAGG 49029 16s
4 GGTTTCCCTTCGGGGCAG GGGAGACAGGGenbank 16s 49029 AB042059
1003 A C C T T C C T T C G G G G C A G - A G G A G A C A G G G enhank 16s 49025 AB042058
1 A C C T IT C C T T C G G G A C A G A G G A C A G G Clostridium elmenteitii
026 GITICCCCTTCGGGGGACAGGGTGACAGGGeobacillus subterraneus 168 AF2763
GGGGCTG - GGGAGACAGG Sulfobacillus disulfidooxidans 168
OZIGILICI ILC GGGGGACACAGGGILACAGGG

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted)  Thursday, September 04, 2003 10:51 AM	Page 19
TGGTGCATGGTTGTCGTCAGCTCGTGTCGT Conservation Conservation Conservations of the Conservation Conse	Consensus #1 Consensus #1 Majority
1090	
1034 TGGTGCATGGTTGTCGTCAGCTCGTGT 43030 1034 TGGTGCATGGTTGTCGTCAGCTGTCGT genbar 1032 TGGTGCATGGTTGTCGTCAGCTCGTGTCGT 49029	43030 16s genbank 16s 43030 AB059664 49029 16s
1032 TGGTGCATGGTTGTCAGCTCGTGTGGT genbar 744	genbank 16s 49029 AB042059
T G G T G C A T G G T T G T C G T C A G C T C G T G T	
100101001001001001001001001001001001001	Clostridium elmenteitii
033 TGGTGCATGGTCGTCAGCTCGTGTCGT	subreitameus us disulfidoox
1047 TGGTGCATGGTTGTCGTCAGCTCGTGTCGT Bacil	Bacillus thermoleovorans ribosomal RNA
GAGATGTTGGGTTAAGTCCCGCAACGAGCG Conser	Consensus #1
GAGATGGGTTAAGTCCGAAAAAAAAAAAAAAAAAAAAAA	Consensus #1
1120 1130 1140	
1064 GAGATGTTCAGTCCCGCAACGAGCG 43030	3030 168
TCAGTCCCGCAACGAGCG	
GAGATGTTGGGTTAAGTCCCGCAACGAGCG	49029 les genbank 16s 49029 AB042059
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	16SRDNA-t7
GAGATGTTGGGTTAAGTCCCGCAACGAGCG	30420
9 GAGATGTTGGGTTAAGTCCCGCAACGAGCG	Clostridium elmenteitii
86 GAGATGTTGGTTAAGTCCCGCAACGAGCG	lbterraneus 16S AF2763
063 G A G A T G T T G G G T T A A G T C C C G C A A C G A G C G	cidans 16S
IGAGATGTTGGGTTAAGTCCCGCAACGICG	Bacillus thermoleovorans ribosomal RNA

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted)  Alignment 2  Thursday, September 04, 2003 10:51 AM
CAACCCTTGA.CTGTTACCAGCACGTConsensus #1
1094 C A A C C C T G G G T T A C C A G C G C G T T G G G G A 23330 168  1094 C A A C C C T T G A C C T G T G T T A C C A G C G C G T T G G G C A 23330 168  1092 C A A C C C T T G A A C T G T T A C C A G C G C A
AGGTGGGACTCACAGGTGACTGCCGGCG- Consensus #1 GGG.A
CTCACAGGTGACTGCCGGCCCTCTCACAGGTGACTGCCGGCCCCTCACAGTTGACTGCCGGCCCCTCACACAGTTGACTGCCGGCCCCCCCC

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted)  Alignment 2  Thursday, September 04, 2003 10:51 AM
TAAGTCGGAGGAAGGCGGGATGACGTCAA Consensus #1
1210 1220 1230
3       TAAGTCGGAGGAAGGCGGGATGACGTCAA       43030 16s         3       TAAGTCGGAGGAAGGCGGGATGACGTCAA       genbank 16s 43030 AB059664         1       TAAGTCGGAGGAAGGCGGGATGACGTCAA       49029 16s         1       TAAGTCGGAGGAAGGCGGGATGACGTCAA       genbank 16s 49029 AB042059         2
1174 AAAAGTCGGAGGAAGGTGGGGATGACGTCAA Geobacillus subterraneus 168 AF276307. 1152 TAAGACGAAGGCGGGGGATGACGTCAA Sulfobacillus disulfidooxidans 168 U349 1165 CAAGTCGGAGGAAGGTGGGGATGACGTCAA Bacillus thermoleovorans ribosomal RNA
ATCATGCCCCTTATGTCCTGGGCTACA Consensus #1
1183 ATCATCATCATGCCCTGGCCTACA genbank 16s 43030 AB059664  1181 ATCATCATGCCTGGCCTACA genbank 16s 43030 AB059664  1181 ATCATCATGCCTTGATGCCTGGGCTACA genbank 16s 49029 AB042059  752

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted) Thursday, September 04, 2003 10:51 AM	Page 22
CACGTGCTACAATGGGCGGTACAA.GGGA. Consensus #1	
1270 1280 1290	
1213 CACGTGCTACAATGGGCGGAACAAAGAAAAGGAAAGGAA	307 U349 RNA
GCGAA. CCGCGAGG. GGAGC. AA. CCCA. A Consensus #1 Consensus #1 GCGAAGCCGCAGGTGGAGCGAACCCCAAA Majority 1300 1310	÷ .
1243 G C G A A G C C G C G A G G C G A A A C C C A A A Genbank 16s 43030 AB059664 1241 G C G A G G C G A G G C G A G C C A A A C C C T G A 49029 16s 1241 G C G A G G C G A G G T G G A G C C A A A C C C T G A 49029 16s 1241 G C G A G A G G T G G A G C A A A C C C T G A genbank 16s 49029 AB042059 1240 G C G A A G G T G G A G C A A A C C C T A A Genbank 16s 49029 AB042059 1240 G C G A A G G T G G A G C A A A C C C A T A Genbank 16s 49025 AB042058 1238 G C G A A G G T G G A G C A A T C C C A T A Genbank 16s 49025 AB042058 1254 G C G A A G G A G G G G A G C A A T C C C A T A Geobacillus subterraneus 16S AF2765 1255 G C G A A C C G C G A G G G G A A T C C C A A A Geobacillus disulfidooxidans 16S 1255 G C G A A C C G C G A G G G G A A T C C C A A A Bacillus thermoleovorans ribosomal	-ed AF276307 s 16S U349

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted)  Thursday, September 04, 2003 10:51 AM	GCCG.TCGTAGTTCGGATTGCAGGCTGC Consensus #1TT	1330 1340 1350	G C C G C T C G T A G T T C G G A TT G C A G G C T G C G enbank 16s 43030 AB059664  G C C G C T C G T A G T T C G G A TT G C A G G C T G C G enbank 16s 49029 AB042059  G C C G T T C G T A G T T C G G A TT G C A G G C T G C G enbank 16s 49025 AB042059  TM A M A C C 4902516SRDNA-t7p_C02_006-1-ed  G C C G T T C G T A G T T C G G A TT G C G A G C T G C G enbank 16s 49025 AB042058  TM A M A C C G C T G C G C T G C G C T G C G C T G C G C	CTCGCCTGCATGAAGCCGGAATTGCTAG Consensus #1	0 0 0 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Alignment Report of Ali 16 Thursday, September 04,			0 0 0 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	υ · υ υ · υ	

Alignment Report of Ali 16S alignement meg ClustalV (Weighted)  Thursday, September 04, 2003 10:51 AM	Page 24
TAATCGCGGATCAGCATGCCGGTGAAT. Consensus #1	
TAATCGCGGATCAGCATGCCGGTGAATC Majority	-
1390 1400 1410	
CGCGGATCAGCA	
CGCGGATCAGCATGCCGGGTGAATA genbank 16s 43030	AB059664
331 TAATCGCGGATCAGCATGCCGCGGTGAATC	
331 TAATCGCGGATCAGCATGCCGCGGTGAATC genbank 16s 49029	AB042059
65	)2 006-1-ed
330 TAATCGCGGATCAGCATGCCGGGTGAATC genbank 16s 49025	AB042058
328 TAATCGCGAATCAGCATGTCGCGGTGAATG	·-
354 TAATCGCGGATCAGCA	us 16S AF276307
332 TAATCGCGGATCAGCATGCCGCGGTGAATC Sulfobacillus	
GCGGATCAGCATGCCGGTGAATA Bacillus therm	ribosomal
C C E	
TICCCGGGCCT	
1420 1430 1440	
G T T T T T T T B	
363 CG T T COTTGTACACCCCCTC genbank 16s 43030	AB059664
361 CG TT CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	
61 C G W T M G G G C C T T G T A C A C C G C C G T C Genbank 16s 49029	AB042059
65 TO C C C - 4902516SRDNA-t7p	_C02_006-1-ed
360 CG T CONTROLACACACCGCCGTC	12058
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384 C G T T CONTROL G G G C C T T G T A	us 16S AF276307
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Alignment 2 Page 25	Consensus #1	Consensus #1	Majority	0	43030 16s	genbank 16s 43030 AB059664	49029 16s	genbank 16s 49029 AB042059	cc-4902516SRDNA-t7p_C02_006-1-ed	genbank 16s 49025 AB042058	Clostridium elmenteitii	Geobacillus subterraneus 16S AF276307	Sulfobacillus disulfidooxidans 168 U349	Bacillus thermoleovorans ribosomal RNA
Alignment Report of Ali 16S alignement.meg ClustalV (Weighted) Thursday, September 04, 2003 10:51 AM	ACACCACGAGAGTCGGCAACCCCGAAGTC Consensus		ACACCACGAGAGTCGGCAACACCGGAAGTC Majority	1450 1460 1470	1393 ACACCACGAGAGTCGGCAACACCCGAAGTC 43030 168	1393 АСАССАС В В В В ТСВ В САРССС В Р В ТС	1391 АСАССАС В В В В Т С В В СА В С С С В А В Т С В	1391 АСАССАСВАВТСВВСААСАССВААВТС	770	1390 ACACCACGAGAGTCGGCAACACCCGAAGTC	1388 АСАССАС 6 ССАТС 6 САДА СТСС САДА ССССВАЯ 6 ССС	1414 ACACCACGAGAG <mark>CTTG</mark> CAACACCGGAAGTC	1392 ACACCACGAGAG <u>TCGA</u> CAACACCCGAAG <u>TC</u>	1405 ACACGAGAGCTCGCAACACCC

WO 2004/063699

Fighted   Alignment 2   Fage 26     A T T G G G T G A A G Consensus #1     A T T G G G T G A A G   Majority   Majority     1520		•	WU 2004/063699	50.104	PC1/US2003/038399
Fighted   Alignment   Alignment   Alignment	Page Zb	Page 26		ed F2763 168 Omal	4 9 06-1-ed 8 16S AF276307 idans 16S U349 ribosomal RNA
Alignment Report of Ali 16S alignement.meg Clusta  Thursday, September 04, 2003 10:51 AM  C C G A A G G T G G G T C G A  1453 C C G A A G G T G G G T C G A  1450 C C G A A G G T G G G T C G A  1450 C C G A A G G T G G G T C G A  1450 C C G A A G G T G G G T C G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T T G A  1450 C C G A A G G T G G G T A G C C  1483 T C G T A A C A A G G T A G C C  1483 T C G T A A C A A G G T A G C C  1480 T C G T A A C A A G G T A G C C  1480 T C G T A A C A A G G T A G C C  1480 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C G T A A C A A G G T A G C C  1470 T C C G A A G C C C  1470 T C C C A A G C C C C C  1470 T C C C A A G C C C C C C  1470 T C C C A A G C T A C C C  1470 T C C C A A G C T A C C C C  1470 T C C C A A G C T A C C C C  1470 T C C C C A A C C C C C C C C C C C  1470 T C C C	Report of All 16S alignement.meg Ciustaly (weignted) September 04, 2003 10:51 AM	Report of Ali 16S alignement.meg ClustalV (Weighted)	day, September 04, 2003 10:51 AM         c c g A A G G T G G G G A T G A T T G G G T G A G Consensus #1         c c g A A G G T G G G T C G A T G A T T G G G T G A A G Majority         1510       1520         C C G A A G G T G G T C G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G A T G A T T G G G T G A A G G T G A T G A T T G G G T G A A G G T G A T G A T T G G G T G A A G G T G G G T G A T G A T T G G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G A G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A T G A T T G G G T G A A G G T G A A G G T G A A G G T G A A G G T G A A G G T G A A G G T G A A G G T G A T G A T T G G G G	CCGAAGGTGGGTTGATGATTGGGTGAAG  CCGAAGGTGAAGCCGATAACTTGGGGTGAAG  CCGAAGGTGAAGCCGATTAACTTGGGGTGAAG  CCGAAGGTGAAGTGATTGGGGTGAAG  CCGAAGGTAGCGATGATTGGGGTGAAG  CCGAAGGTAGCCGTXXXXXXXXXXXXXXXXXXXXXXXXXX	1540

Alignment Report of Ali 16S alignement.meg ClustalV (Weighted) Thursday, September 04, 2003 10:51 AM	Alignment 2 Page 27 O
	Consensus #1
	Consensus #1
XXXXXXXXXXXXXXXX	Majority
1570	
1500	43030 168
1513 G G T T G G A T	genbank 16s 43030 AB059664
1497	49029 16s
1510 G G T T G G A T	genbank 16s 49029 AB042059
770	cc-4902516SRDNA-t7p_C02_006-1-ed
1508 GGTTGGA	genbank 16s 49025 AB042058
1492	Clostridium elmenteitii
1533 GGCTGGATCACCTCCT	Geobacillus subterraneus 16S AF276307
1496	Sulfobacillus disulfidooxidans 16S U345
1428	Bacillus thermoleovorans ribosomal RNA

of the Consensus, otherwise show '.'.

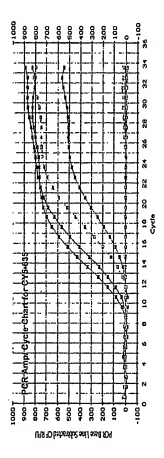
Consensus 'Consensus #1': When 60% (6) match the residue of the Consensus show the residue

Consensus 'Consensus #1': When all match the residue of the Consensus show the residue of the Consensus, otherwise show '.'.

at 40% fill) residues that match the Decoration 'Decoration #1': Shade (with black consensus named 'Consensus #1' exactly.

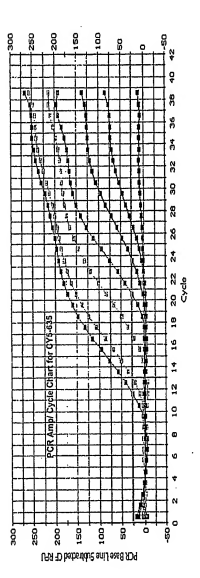
Decoration 'Decoration #1': Box residues that match the Consensus exactly





In addition, G. stearothermophilus can be seen in green. Other curves below the base line include the blank control (red) and the tests for Lactococcus lactis C2 (it blue), P. putida 49L/51 (purple) and E. coli DH5 peranggusing the same primer and probe set.

Figure 10

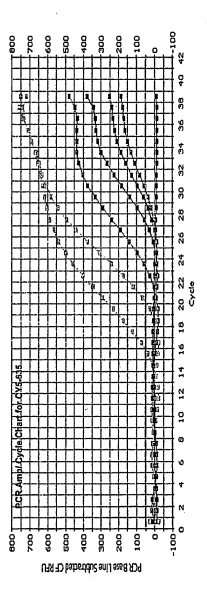


. Dilutions 10⁰ (red), 10⁻¹ (orange), 10⁻² (it. bitte), 10⁻³ (purple), 10⁻⁴ (black), and 10⁻⁵ (blue) can be seen above the baseline. The blank control (brown) is below the baseline. 15

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Dilutions 10⁻¹ (orange), 10⁻² (h. hhuc), 10⁻³ (purple), 10⁻⁴ (black), 10⁻⁵ (blue) and 10⁻⁶ (green) can be seen above the baseline. The blank control (red) is below the baseline.

# Rights 11

GTTGATTACGTCCCTGCCCTTTGTACACACACGCGAATTCCAGCACACTGGCGGCCGTTACTAGTGGATCCGAGCTCGG AAAAGGCCGCGTGCTGGCGTTTTINCNTANGCTCGCCCCCTGACAGCATNCAAAATCGACGCTCAGTCNNANGTGGCGAAC TTTTCCACNTTCTTAGAGGGACTATCGGTTTTCAAGCCGATGGAAGTTTGAGGCAATAACAGGTCTGTGATGCCCTTAGA CTCCGTCGTGCTGGGGATAGAGCATTGTAATTATTGCTCTTCAACGAGGAATTCCTAGTAAGCGCAAGTCATCAACTTGC TACCAAGCTTGGCGTAATCATGGTCATAGCTGTTTTCCTGTGTGAAATTGTTATCCGCTCACAATTCCACACATACGA ITTCCAGTCGGGAAACCTGTGCTGCCAGCTGCATTAATGAATCGGCCAACGCGGGGAGAGAGGCGGTTTGCGTATTGGGC STAATACGGTTATCCACAGAATCAGGGGATAACGCAGGAAAGAACATGTGAGCAAAAAGGCCAGCANANGCCAGGANCCGT CCGNNGGANATAAGATACNNGCGTTNCCCCTGNANCTCCNCNTGGCTNTCNGNTCNANCNGNCGNTANGGAANCTGNCNČ CGTTCTGGGCCGCACGCGCGCTACACTGACGGAGCCAGCGAGTCTAACCTTGGCCGAGAGGTCTGGGTAATCTTGTGAAA ATTGGGCCCTCTANAGCATGCTCGACGGCCGCCAGTGTGATGGATATCTGCAGAATTCGGCTTTGCATGGCCGTTCTTAG Zygosaccaromyces S 10

15

# Figure 13

AGGTNNGANCAGGAGAGCGCACGAGGGAGCTNCAGGGGGAAACGCCTGGGGATCTTNATAGTCCNGTCGGGTTCNCCACNT CTATCGGCTCAAGCCGATGGAAGTGCGCGGCGATAACAGGTCTGTGATGCCCTTAGATGTTCTGGGCCGCACGCGCGCTA GANGNCNNCCCNNANTNNATCCTNAGCNGAGTNGNNAAGCGCNCGTTNCCGANGGAGAAGNGGACAGGTNTCCGTANCGC GTTGTGTGGAATTGTGAGCGGATAACAATTTTCACACAGGAAACAGCTATGACCATGATTACGCCAAGGTTTGGTACCGAGC GCTTAALTIGCGATAACGAACGAGGACCTCGGCCCTTAAATAGCCCCGGTCCGCATTTGCGGGCCGCTGGCTTCTTAAGGGGA CACTGACAGGGCCAGCGAGTACATCACCTTAACCGAGAGGTTTGGGTAATCTTGTTAAACCCTGTCGTGCTGGGGATAGA TGAGTGAGCTGATACCGCTCGCCGCAGCCGAACGACCGAGCGCAGCGAGTCAGTGAGCGAGGGAAGCGGAAGAGCGCCCAA AGTGAGCGCAACGCAATTAATGTGAGTTAGCTCACTCATTAGGCACCCCAGGCTTTACACTTTATGCTTCCGGCTCGTAT <u> TTGACTTGAGCGTCGATTTTTGTGATGCTCGTCAGGGGGCGGAGCNTATGGAAAACGCCAGCAACGCGGCCTTTTTACGGTT</u> CTGGCNTTTGCTGGCCTTTTTGCTCACATGTTCTTTCCTGCGTTATCCCCTGATTCTGTGGATAACCGTATTACCGCCTT TACGCAAACCGCCTCTCCCCGCGCGTTGGCCGATTCATTAATGCAGCTGGCACGACAGGTTTCCCGACTGGAAAGCGGGC TCGGATCCACTAGTAACGGCCGCCAGTGTGCTGGAATTCGGCTTTTGCATGGCCGTTCTTAGTTGGTGGAGTGATTTGTCT 3CATTGCAATTATTGCTCTTCAACGAGGAATGCCTAGTAGGCACGAGTCATCAGCTCGTGCCGATTACGTCCCTGCCCTT TGTACACACAAGCCGAATTCTGCAGATATCCATCACACTGGCGGCCGTCGAGCATGCTNTAGAGGGCCCAAT Penecillium digitatum 10 S 15

# Figure 14

# Byssochlamys fulva

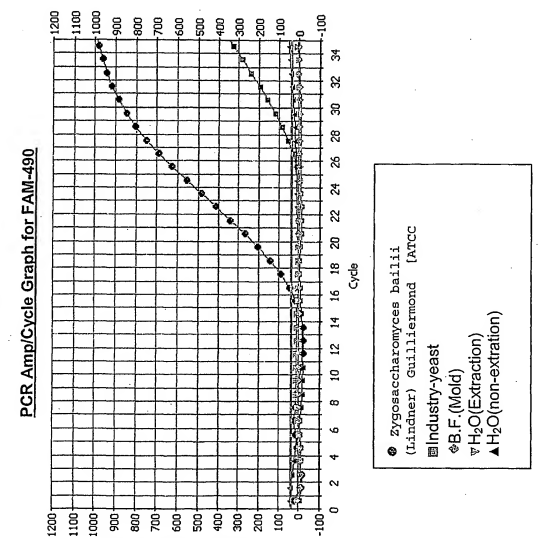
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NNNNNNNNNNNTNTCNNGGGNNNGNGCNNNNGNAAANNNCCNGCANNNNGCCNTTNNNNGNTNNNGGCCCNTNNGHNGHC NININGNI CACANGI INNI CONNGCGNINI CCCNNGNI INNIGNGGA TAACNGIA L'INCCGCCINNGAGIGAGNI GA IACCGC AATGTGAGTTAGCTCACTCATTAGGCACCCCAGGCTTTACACTTTATGCTTCCGGCTCGTATGTTGTGTGGAATTGTGAG GGATAACAATTTCACACAGGAAACAGCTATGACCATGATTACGCCAAGCTTGGTACCGAGCTCGGATCCACTAGTAACG SCCGCCAGTGTGCTGGAATTCGGCTTTGCATGGCCGTTCTTAGTTGGTGGAGTGATTTGTCTGCTTAATTGCGATAACGA <u> ACGAGACCTCGGCTCTTAAATAGCCCGGTCCGCGTTTGCGGGCCGCTGGCTTCTTAGGGGGACTATCGGCTCAAGCCGAT</u> STACATCACCTTGGCCGAGAGGTCTGGGTAATCTTGTTAAACCCTGTCGTGGGGGATAGAGCATTGCAATTATTGCTC ITCAACGAGGAATGCCTAGTAGGCACGAGTCATCAGCTCGTGCCGATTACGTCCCTGCCCTTTGTACACAAGCCGAAT CCGCGCGTTGGCCGATTCATTAATGCAGCTGGCACGACAGGTTTCCCGACTGGAAAGCGGGCAGTGAGCGCAATTT <u> PCTGCAGATATCCATCACACTGGCGGCCGTCGAGCATGCTNTAGAGGGCCCCAAT</u>

15





PCR Base Line Subtracted CF RFU

Figure 16

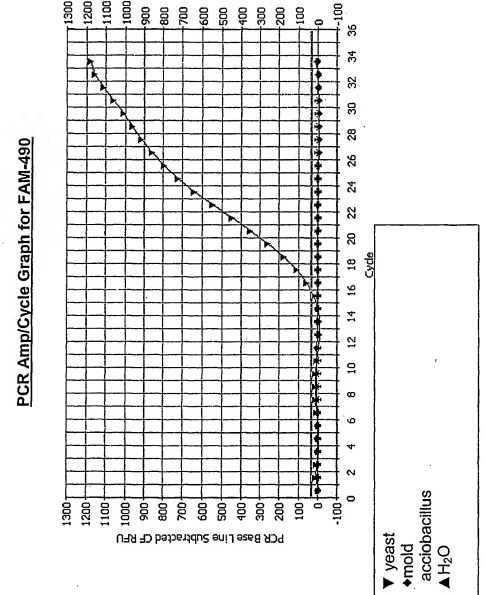
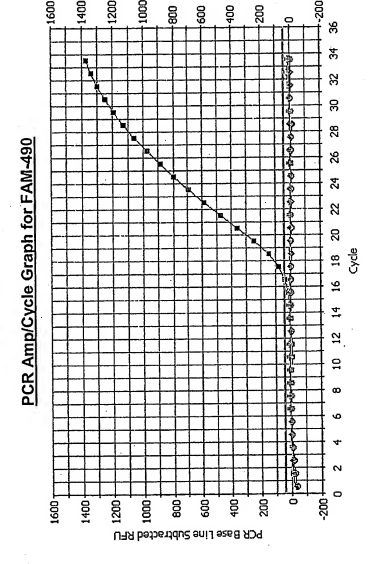


Figure 17





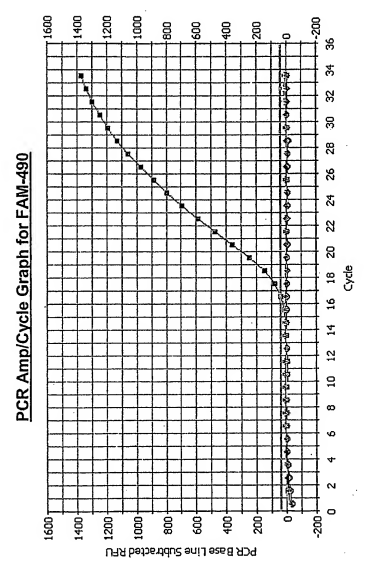


Figure 19

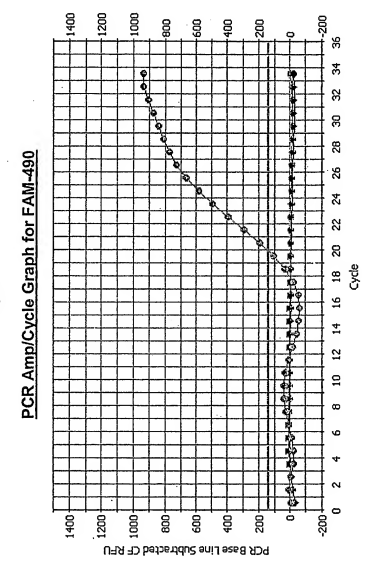
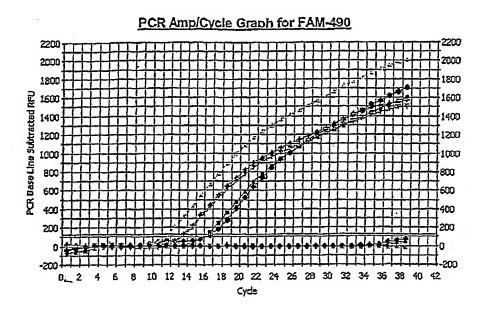
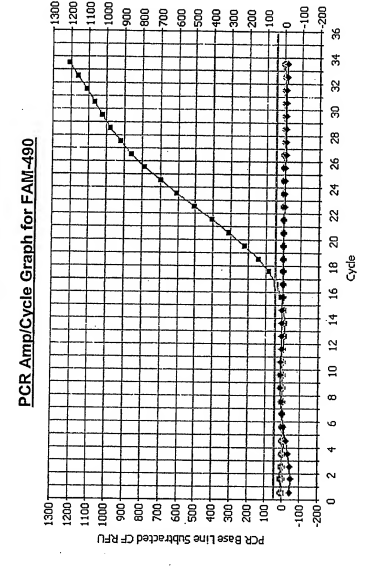


Figure 20





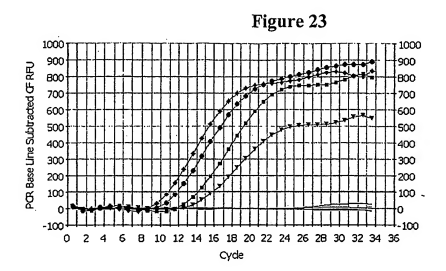


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# Figure 22

```
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGC|TGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CCCAGTTCGGATTGAGGGCTGCAACTCGCCCCCATGAAGT
CTCAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGTAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CTCAGTTCGGATTGCAGGCTGCAACTCGCCTGCATGAAGC
CGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATACGTTC
CGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATACGTTC
CGGAATTGCTAGTAATCGCBGATCAGCATGCCGCGGTGAATCCGTTC
CGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATCCGTTC
CGGAATTGCTAGTAATCGCBGATCAGCATGCCGCGGTGAATCCGTTC
CGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATCCGTTC
TGGAGTTGCTAGTAATCGCGAATCAGCAT GTCGCGGTGAATGCGTTC
CGGAATCGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATACGTTC
CGGAATTGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATCCGTTC
CGGAATCGCTAGTAATCGCGGATCAGCATGCCGCGGTGAATACGTTC
CCGGGCCTTGTACACACCGCCCGTCACACCGCGAGAGTCGGCAACAC
CCGGGCCTTGTACACACCGCCCGTCACACCCGAGAGTCGGCAACAC
CCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGTCGGCAACAC
CCGGGCCTTGTACACACCGCCCGTCACAQCACGAGAGTCGGCAACAC
CCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGTCGGCAACAC
CCGGGCCTTGTACACACCGCCGTCACACCACGAGAGTCGGCAACAC
CCGGGTCTTGTACACACCGCCCGTCACACCCACGGAAGTCGGAAGCAC
CCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGCTTGCAACAC
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CCGGGCCTTGTACACACCGCCCGTCACACCACGAGAGCTCGCAACAC
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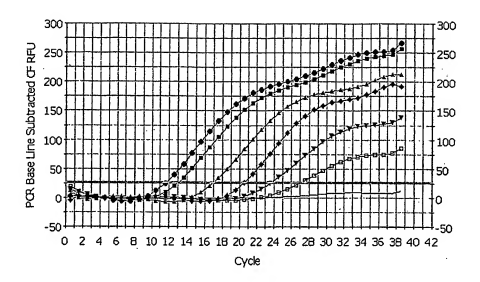
a 16s rDNA Sequences in the alignment are 16S rDNA sequences from the following organisms (GenBank accession numbers follow if applicable): 1) A. acidocaldarius strain ATCC 43030, 2) A. acidocaldarius strain DSM 454 (AB059664), 3) A. cycloheptanicus strain ATCC 49029, 4) A. cycloheptanicus strain DSM 4006 (AB042059), 5) A. acidoterrestris strain ATCC 49025, 6) A. acidoterrestris strain DSM 3923 (AB042058), 7) Clostridium elmenteitii isolate E2SE1-B (AJ271453), 8) Geobacillus subterraneus strain K (AF276307), 9) Sulfobacillus disulfidooxidans SD-11 (U34974), and 10) B. thermoleovorans strain ATCC 43513 (M77488) Note CC16S-R is in 5' to 3' orientation in alignment. Actual primer sequence is the reverse complement.



In addition, G. stearothermophilus ATCC 10149 (▼) can be detected. Curves below the base line include the blank control, Lactococcus lactis C2, P. putida 49L/51, and E. coli DH5α. This is a representative curve of repeated trials.

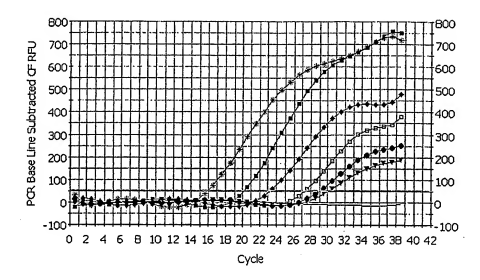
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Figure 24



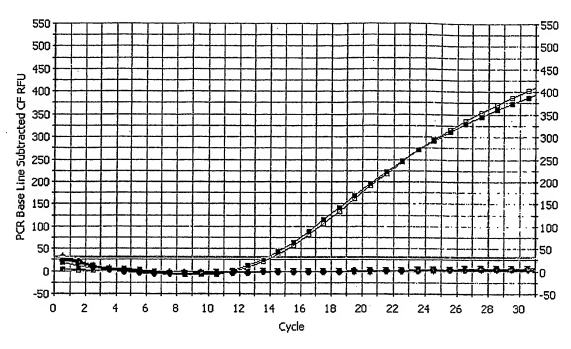
. Ten-fold serial dilutions were performed, and cell numbers^a represented at each curve are as follows: 1.6x10⁷ CFU/ml (•), 1.6x10⁶ CFU/ml (■), 1.6x10⁵ CFU/ml (▲), 1.6x10⁴ CFU/ml (♦), 1.6x10³ CFU/ml (▼), and 1.6x10² CFU/ml (□) appear above the baseline. The water control is below the baseline. This is a representative curve of repeated trials. ^a Cell numbers were calculated by finding the CFU/ml of plated samples and multiplying by the dilution level of the representative curve.

Figure 25



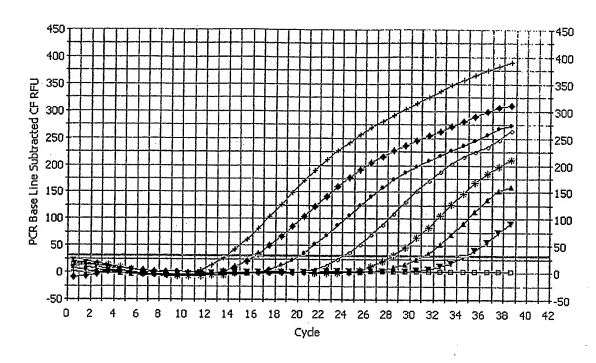
Cell counts at each curve are as follows: 6.3x10⁶ CFU/ml (*), 6.3x10⁵ CFU/ml (■), 6.3x10⁴ CFU/ml (◆), 6.3x10³ CFU/ml; (□), 6.3x10² CFU/ml (•), and 6.3x10¹ CFU/ml (▼) appear above the baseline. The water control is below the baseline. This is a representative curve of repeated trials.

Figure 26



A. acidoterrestris ATCC 49025 (■), A. acidocaldarius ATCC 43030 (□), 6 other bacteria under the detection baseline: Bacillus subtilis OSU 456, Psudomonas putida 49L/51, Ecoli DH5α, Listeria monocytogenes V7, Lactococcus lactis ML3 and Geobacillus ATCC 10149 ■ Alicyclobacillus acidoterrestris ATCC49025; □ Alicyclobacillus acidocaldarius ATCC43030; + Bacillus subtilis OSU456; • E.coli DH5α; ▲ Psudomonas putida 49L/51; ▼ Geobacillus ATCC 10149; ◆ Lactococcus lactis ML3; ○Listeria monocytogenes V7

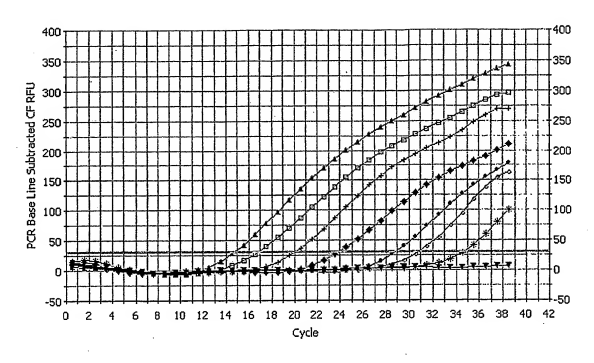
Figure 27



- 5 ▲ 1ml saline with 4× 10⁶ CFU/ml Alicyclobacillus acidoterrestris ATCC49025
  - $\ \square$  1ml saline with 4×  $10^5\, \text{CFU/ml}$  Alicyclobacillus acidoterrestris ATCC49025
  - + 1ml saline with 4× 10⁴ CFU/ml Alicyclobacillus acidoterrestris ATCC49025
  - ♦ 1ml saline with 4×10³ CFU/ml Alicyclobacillus acidoterrestris ATCC49025
  - 1ml saline with 4× 10² CFU/ml *Alicyclobacillus acidoterrestris* ATCC49025
- 10  $\circ$  1ml saline with 4× 10¹ CFU/ml *Alicyclobacillus acidoterrestris* ATCC49025
  - *1ml saline with 4CFU/ml Alicyclobacillus acidoterrestris ATCC49025

▼ 1ml saline

Figure 28



5

15 ▼1ml apple Juice

# (19) World Intellectual Property Organization

International Bureau





# (43) International Publication Date 29 July 2004 (29.07.2004)

**PCT** 

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60/430,202 2 De

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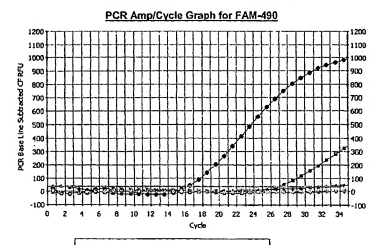
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- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (regional): ARIPO patent (BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE,

[Continued on next page]

(54) Title: RAPID DETECTION OF MICROORGANISMS



- Zygosaccharomyces bailii (Lindner) Guilliermond (ATCC
- Industry-yeast
- ⇔B.F.(Mold)
- ▼H₂O(Extraction)
- ▲H₂O(non-extration)

(57) Abstract: Tools and methods for detecting the presence bacteria, yeast and mold in a sample obtained from a food sample are provided. The methods employ a polymerase chain reaction and primer and probe sets that are based on the 16S rRNA and squalene-hopene cyclase genes of Alicyclobacillus and Geobacillus and the 18S rDNA gene of mold and yeast. The present invention also relates to primer and probe sets. Each primer and probe set comprises a forward primer and a reverse primer, both of which are from 15 to 35 nucleotides in length and a probe.

## WO 2004/063699 A3



SI, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

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- of inventorship (Rule 4.17(iv)) for US only

#### Published:

- with international search report

- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments
- (88) Date of publication of the international search report:
  16 December 2004

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# INTERNATIONAL SEARCH REPORT

International application No.

PCT/US03/38399

A. CLASSIFICATION OF SUBJECT MATTER					
IPC(7) : C12Q 1/68; C12P 19/34; C07H 21/04 US CL : 435/6, 91.2: 536/23.7. 24.32, 24.33					
US CL : 435/6, 91.2; 536/23.7; 24.32, 24.33 According to International Patent Classification (PC) or to both national classification and IPC					
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U.S. :	435/6, 91.2; 536/23.7, 24.32, 24.33	ed by classification symbols)			
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·X	WO 02/070728 A2 (THE JOHNS HOPKINS UNI	VERSITY) 12 September 2002	25-29		
Α	(12.09.2002), see entire reference, particularly pa YAMAZAKI, K. et al. Specific primers for detec by RT-PCR. Letters in Applied Microbiology. N see entire reference.	tion of Alicyclobacillus acidoterrestris	1, 5, 9, 13, 17, 21-23		
	see entire reference.				
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International application No.

PCT/US03/38399

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2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
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